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(57) Abstract

The present invention provides recombinant DNA comprising a transcription promoter and a downstream sequence to be expressed, in operable linkage therewith, wherein the transcription promoter comprises a region found upstream of the open reading frame of a highly expressed *Phaffia* gene, preferably a glycolytic pathway gene, more preferably the gene coding for Glyceraldehyde-3-Phosphate Dehydrogenase. Further preferred recombinant DNAs according to the invention contain promoters of ribosomal protein encoding genes, more preferably wherein the transcription promoter comprises a region found upstream of the open reading frame encoding a protein as represented by one of the amino acid sequences depicted in any one of SEQIDNOs: 24 to 50. According to a further aspect of the invention an isolated DNA sequence coding for an enzyme involved in the carotenoid biosynthetic pathway of *Phaffia rhodozyma* is provided, preferably wherein said enzyme has an activity selected from isopentenyl pyrophosphate isomerase activity, geranylgeranyl pyrophosphate synthase activity, phytoene synthase activity, phytoene desaturase activity and lycopene cyclase activity, still more preferably those coding for an enzyme having an amino acid sequence selected from the one represented by SEQIDNO: 13, SEQIDNO: 15, SEQIDNO: 17, SEQIDNO: 21 or SEQIDNO: 23. Further embodiments concern vectors, transformed host organisms, methods for making proteins and/or carotenoids, such as astaxanthin, and methods for isolating highly expressed promoters from *Phaffia*.

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Improved methods for transforming *Phaffia* strains, transformed *Phaffia* strains so obtained and recombinant DNA in said methods

Technical field

The present invention relates to methods for transforming *Phaffia* yeast, transformed *Phaffia* strains, as well as recombinant DNA for use therein.

Background of the invention

Methods for transforming the yeast *Phaffia rhodozyma* have been disclosed in European patent application 0 590 707 A1. These methods involve incubation of protoplasts with DNA or incubation of *Phaffia* cells with DNA followed by lithium acetate treatment. The recombinant DNA used to transform *Phaffia* strains with either of these methods comprised a *Phaffia* actin gene promoter to drive expression of the selectable marker genes coding for resistance against G418 or phleomycin. The methods involve long PFG and lithium acetate incubation times and transformation frequencies are low. When protoplasts are used, the transformation frequency is dependent on the quality of the protoplast suspension, making the procedure less reliable.

Recently a method for transforming *Phaffia* strains has been reported by Adrio J.L. and Veiga M.(July 1995, Biotechnology Techniques Vol. 9, No. 7, pp. 509-512). With this method the transformation frequencies are in the range of 3 to 13 transformants per µg DNA, which is low. A further disadvantage of the method disclosed by these authors consists in increased doubling time of the transformed cells. The authors hypothesised that this may be due to interference of the autonomously replicating vector with chromosome replication.

Clearly, there is still a need for a reliable and efficient method of transforming *Phaffia* strains with foreign DNA. It is an objective of the present invention to provide methods and means to achieve this. It is a further objective of the invention to optimize expression of certain genes in *Phaffia* rhodozyma in order to make *Phaffia* a more suitable production host for certain valuable compounds.

Summary of the invention

The invention provides a method for obtaining a transformed *Phaffia* strain, comprising the steps of contacting cells or protoplasts of a *Phaffia* strain with recombinant DNA under conditions conducive to uptake thereof, said recombinant DNA comprising a transcription promoter and a downstream sequence to be expressed which is heterologous to said transcription promoter, in operable linkage therewith, identifying *Phaffia* rhodozyma cells or protoplasts having obtained the said recombinant DNA in expressible form, wherein the transcription promoter comprises a region that is found upstream of the open reading frame of a highly expressed *Phaffia* gene. According to a preferred embodiment of the invention said highly expressed *Phaffia* gene is a glycolytic pathway gene, more preferably the glycolytic pathway gene is coding for Glyceraldehyde-3-Phosphate Dehydrogenase

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(GAPDH). According to one aspect of the invention, said heterologous downstream sequence comprises an open reading frame coding for resistance against a selective agent, such as G418 or phleomycin.

Another preferred method according to the invention is one, wherein said recombinant DNA comprises further a transcription terminator downstream from the said DNA to be expressed, in operable linkage therewith, which transcription terminator comprises a region found downstream of the open reading frame of a *Phaffia* gene. It is still further preferred, that the recombinant DNA is in the form of linear DNA.

Another preferred embodiment comprises, in addition to the steps above, the step of providing an electropulse after contacting of *Phaffia* cells or protoplasts with DNA.

According to another embodiment the invention provides a transformed *Phaffia* strain capable of high-level expression of a heterologous DNA sequence, which strain is obtainable by a method according to the invention. Preferably, said *Phaffia* strain contains at least 10 copies of the said recombinant DNA integrated into its genome, such as a chromosome, particularly in the ribosomal DNA locus of said chromosome.

The invention also provides recombinant DNA comprising a transcription promoter and a heterologous downstream sequence to be expressed, in operable linkage therewith, wherein the transcription promoter comprises a region found upstream of the open reading frame of a highly expressed *Phaffia* gene, preferably a glycolytic pathway gene, more preferably a gene coding for Glyceraldehyde-3-Phosphate Dehydrogenase.

Also provided is recombinant DNA according to the invention, wherein the heterologous downstream sequence comprises an open reading frame coding for reduced sensitivity against a selective agent, preferably G418 or phleomycin. Said recombinant DNA preferably comprises further a transcription terminator downstream from the said heterologous DNA sequence to be expressed, in operable linkage therewith.

Further aspects of the invention concern a microorganism harbouring recombinant DNA according to the invention, preferably *Phaffia* strains, more preferably *Phaffia* rhodozyma strains, as well as cultures thereof.

According to still other preferred embodiments isolated DNA fragments are provided comprising a *Phaffia* GAPDH-gene, or a fragment thereof, as well as the use of such a fragment for making a recombinant DNA construct. According to one embodiment of this aspect said fragment is a regulatory region located upstream or downstream of the open reading frame coding for GAPDH, and it is used in conjunction with a heterologous sequence to be expressed under the control thereof.

The invention according to yet another aspect, provides a method for producing a protein or a pigment by culturing a *Phaffia* strain under conditions conducive to the production of said protein or pigment, wherein the *Phaffia* strain is a transformed *Phaffia* strain according to the invention.

According to another aspect of the invention, a method for obtaining a transformed *Phaffia* strain, comprising the steps of

contacting cells or protoplasts of a *Phaffia* strain with recombinant DNA under conditions conducive to uptake thereof,

said recombinant DNA comprising a transcription promoter and a downstream sequence to be expressed in operable linkage therewith,

identifying *Phaffia rhodozyma* cells or protoplasts having obtained the said recombinant DNA in expressible form,

wherein the downstream sequence to be expressed comprises an isolated DNA sequence coding for an enzyme involved in the carotenoid biosynthetic pathway of *Phaffia rhodozyma*. Preferably, said enzyme has an activity selected from geranylgeranyl pyrophosphate synthase (*crt*E), phytoene synthase (*crt*B), phytoene desaturase (*crt*I) and lycopene cyclase (*crt*Y), more preferably an enzyme having an amino acid sequence selected from the one represented by SEQIDNO: 13, SEQIDNO: 15, SEQIDNO: 17 and SEQIDNO: 19. According to a further embodiment, the transcription promoter is heterologous to said isolated DNA sequence, such as a glycolytic pathway gene in *Phaffia*. Especially preferred according to this embodiment is the Glyceraldehyde-3-Phosphate Dehydrogenase (GAPDH) gene promoter.

Also provided is a transformed *Phaffia* strain obtainable by a method according to the invention and capable of expressing, preferably over-expressing the DNA sequence encoding an enzyme involved in the carotenoid biosynthesis pathway gene.

The invention is also embodied in recombinant DNA comprising an isolated DNA sequence according to the invention, preferably in the form of a vector.

Also claimed is the use of such a vector to transform a host, such as a Phaffia strain.

A host obtainable by transformation, optionally of an ancestor, using a method according to any one of claims 1 to 5, wherein said host is preferably capable of over-expressing DNA according to the invention.

According to a further embodiment a method is provided for expressing an enzyme involved in the carotenoid biosynthesis pathway, by culturing a host according to the invention under conditions conducive to the production of said enzyme. Also provided is a method for producing a carotenoid by cultivating a host according to the invention under conditions conducive to the production of carotenoid.

The following figures further illustrate the invention.

Description of the Figures

Fig. 1. Mapping of the restriction sites around the *Phaffia rhodozyma* GAPDH gene. Ethidium bromide stained 0.8 % agarose gel (A) and Southern blot of chromosomal DNA (B) and cosmid pPRGDHcos1 (C) digested with several restriction enzymes and hybridized with the 300-bp PCR fragment of the *Phaffia rhodozyma* GAPDH gene. Lane 1, DNA x Kpnl; 2, xPstl; 3, xSmal; 4, xSphl; L, lambda DNA digested with BstEII; 5, xSstl; 6, xXbal and 7, xXhol.

The blot was hybridized in 6 x SSC, 5 x Denhardt's, 0.1 % SDS, 100 ng/ml herring sperm DNA at 65°C and washed with 0.1 x SSC/0.1% SDS at 65°C. Exposure time of the autoradiogram was 16 h for the cosmid and 48 h from the blot containing the chromosomal DNA.

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Fig. 2. The organisation of two subclones; pPRGDH3 and derivative (A) and pPRGDH6 and derivatives (B) containing (a part of) the GAPDH gene of *Phaffia rhodozyma*. The PCR probe is indicated by a solid box. The direction and extent of the sequence determination is indicated by arrows.

solid boxes: GAPDH coding sequence

open box: 5' upstream and promoter region of GAPDH

open box: 3' non-coding Phaffia rhodozyma GAPDH sequence

solid line: GAPDH intron

hatched box: Poly-linker containing sites for different restriction enzymes

dotted line: deleted fragments

Fig. 3. Cloning diagram of *Phaffia* transformation vector, pPR2.

solid box: 5' upstream and promoter sequence of GAPDH

hatched box: G418 solid line: pUC19

open box: ribosomal DNA of Phaffia rhodozyma

Only restriction sites used for cloning are indicated.

Fig 4 Construction of pPR2T from pPR2T.

Solid box (BamHI - HindIII fragment): GAPDH transcription terminator from Phaffia.

All other boxes and lines are as in Fig. 3. Only relevant details have been depicted.

- Detailed physical map of pGB-Ph9. bps = basepairs; rDNA ribosomal DNA locus of *Phaffia*; act.pro 2 = actin transcription promoter; act.1 5' non-translated and aminoterminal region of the open reading frame; NON COD. = non-coding region downstream of G418-gene;
 - Fig. 6. Detailed physical map of pPR2. GPDHpro = GAPDH transcription promoter region from *Phaffia*. Other acronyms as in Fig. 5.
- Fig. 7. Detailed physical map of pPR2T. Tgdh = GAPDH transcription terminator of *Phaffia*. All other acronyms as in Fig. 5 and 6.
 - Fig. 8. Overview of the carotenoid biosynthetic pathway of Erwinia uredovora.
 - Fig. 9. Representation of cDNA fragments and a restriction enzyme map of the plasmids pPRcrtE (A); pPRcrtB (B), pPRcrtI (C) and pPRcrtY (B).

Detailed description of the invention

The invention provides in generalised terms a method for obtaining a transformed *Phaffia* strain, comprising the steps of

contacting cells or protoplasts of a *Phaffia* strain with recombinant DNA under conditions conducive to uptake thereof,

said recombinant DNA comprising a transcription promoter and a downstream sequence to be expressed which is heterologous to said transcription promoter, in operable linkage therewith,

identifying *Phaffia rhodozyma* cells or protoplasts having obtained the said recombinant DNA in expressible form,

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wherein the transcription promoter comprises a region that is found upstream of the open reading frame of a highly expressed *Phaffia* gene.

In order to illustrate the various ways of practicing the invention, some embodiments will be high-lighted and the meaning or scope of certain phrases will be elucidated.

The meaning of the expression recombinant DNA is well known in the art of genetic modification, meaning that a DNA molecule is provided, single or double stranded, either linear or circular, nicked or otherwise, characterised by the joining of at least two fragments of different origin. Such joining is usually, but not necessarily done *in vitro*. Thus, within the ambit of the claim are molecules which comprise DNA from different organisms or different genes of the same organism, or even different regions of the same gene, provided the regions are not adjacent in nature. The recombinant DNA according to the invention is characterised by a transcription promoter found upstream of an open reading frame of a highly expressed *Phaffia* gene, fused to a heterologous DNA sequence. With heterologous is meant 'not naturally adjacent'. Thus the heterologous DNA sequence may be from a different organisms, a different gene from the same organism, or even of the same gene as the promoter, provided that the downstream sequence has been modified, usually *in vitro*. Such modification may be an insertion, deletion or substitution, affecting the encoded protein and/or its entrance into the secretory pathway, and/or its post-translational processing, and/or its codon usage.

The strong transcription promoter according to the invention must be in operable linkage with the heterologous downstream sequence in order to allow the transcriptional and translational machinery to recognise the starting signals. The regions upstream of open reading frames of highly expressed Phaffia genes contain TATA-like structures which are positioned at 26 to about 40 nucleotides upstream of the cap-site; the latter roughly corresponds with the transcriptional start site. Thus in order to allow transcription of the heterologous downstream sequence to start at the right location similar distances are to be respected. It is common knowledge, however, that there is a certain tolerance in the location of the TATA-signal relative to the transcription start site. Typically, mRNAs of the eukaryotic type contain a 5'-untranslated leader sequence (5'-utl), which is the region spanning the transcription start site to the start of translation; this region may vary from 30 to more than 200 nucleotides. Neither the length nor the origin of the 5'-utl is very critical; preferably it will be between 30 and 200 nucleotides. It may be from the same gene as the promoter, or it may be from the gene coding for the heterologous protein. It is well known that eukaryotic genes contain signals for the termination of transcription and/or polyadenylation, downstream of the open reading frame. The location of the termination signal is variable, but will typically be between 10 and 200 nucleotides downstream from the translational stop site (the end of the open reading frame), more usually between 30 and 100 nucleotides downstream from the translational stop site. Although the choice of the transcription terminator is not critical, it is found, that the when the terminator is selected from a region downstream of a Phaffia gene, preferably of a highly expressed Phaffia gene, more preferably from the GAPDH-encoding gene, the level of expression, as well as the frequency of transformation is improved.

It was found that significant numbers of clones were obtained which could grow on very high G418 concentrations (up to, and over, 1 mg/ml). Transcription promoters according to the invention are

said to be from highly expressed genes, when they can serve to allow growth of transformed Phaffia cells, when linked to a G418 resistance gene as disclosed in the Examples, in the presence of at least 200 µg/ml, preferably more than 400, even more preferably more than 600, still more preferably more than 800 µg/ml of G418 in the growth medium. Examples of such promoters are, in addition to the promoter upstream from the GAPDH-gene in Phaffia, the promoters from Phaffia genes which are homologous to highly expressed genes from other yeasts, such as Pichia, Saccharomyces, Kluyveromyces, or fungi, such as Trichoderma, Aspergillus, and the like. Promoters which fulfill the requirements according to the invention, may be isolated from genomic DNA using molecular biological techniques which are, as such, all available to the person skilled in the art. The present invention provides a novel strategy for isolating strong promoters from Phaffia as follows. A cDNA-library is made from Phaffia mRNA, using known methods. Then for a number of clones with a cDNA insert, the DNA fragment (which represents the cDNA complement of the expressed mRNA) is sequenced. As a rule all fragments represent expressed genes from Phaffia. Moreover, genes that are abundantly expressed (such as the glycolytic promoters) are overrepresented in the mRNA population. Thus, the number of DNA-fragments to be sequenced in order to find a highly expressed gene, is limited to less than 100, probably even less than 50. The sequencing as such is routine, and should not take more than a couple of weeks. The nucleotide sequences obtained from this limited number of fragments, is subsequently compared to the known sequences stored in electronic databases such as EMBL or Geneseq. If a fragment shows homology of more than 50% over a given length (preferably more than 100 basepairs) the fragment is likely to represent the Phaffia equivalent of the gene found in the electronic database. In yeasts other than Phaffia, a number of highly expressed genes have been identified. These genes include the glycolytic phosphoglucoisomerase, phosphofructokinase, phosphotrioseisomerase, genes, phosphoglucomutase, enolase, pyruvate kinase, alcohol dehydrogenase genes (EP 120 551, EP 0 164 556; Rosenberg S. et al., 1990, Meth. Enzymol.: 185, 341-351; Tuite M.F. 1982, EMBO J. 1, 603-608; Price V. et al., 1990, Meth. Enzymol. 185, 308-318) and the galactose regulon (Johnston, S.A. et al., 1987, Cell 50, 143-146). Accordingly, those Phaffia cDNA fragments that are significantly homologous to the highly expressed yeast genes (more than 40%, preferably more than 50% identity in a best match comparison over a range of more than 50, preferably more than 100 nucleotides) should be used to screen a genomic library from Phaffia, to find the corresponding gene. Employing this method, 14 higly expressed mRNAs from Phaffia rhodozyma have been copied into DNA, sequenced, and their (putative) open reading frames compared to a nucleic acid and amino amino acid sequence databases. It turned out that 13 out of these fourteen cDNAs coded for ribosomal protein genes, of which one coded simultaneously to ubiquitin; one cDNA codes for a glucose-repressed gene. The isolation of the genes and the promoters usually found upstream of the coding regions of these genes is now underway, and it is anticipated that each of these transcription promoters may advantageously be used to express heterologous genes, such as carotenoid biosynthesis genes. Among the genes and transcription promoters especially preferred according to this invention are the promoter found upstream of the ubiquitinribosomal 40S protein corresponding to the cDNA represented in SEQIDNO:10, the glucose-repressed cDNA represented in SEQIDNO:26, the 40S ribosomal protein S27 encoding cDNA represented in

SEQIDNO:28, the 60S ribosomal protein P1α encoding cDNA represented by SEQIDNO:30, the 60S ribosomal protein L37e encoding cDNA represented in SEQIDNO:32, the 60S ribosomal protein L27a encoding cDNA represented in SEQIDNO:34, the 60S ribosomal protein L25 encoding cDNA represented in SEQIDNO:36, the 60S ribosomal protein P2 encoding cDNA represented in SEQIDNO:38, the 40S ribosomal protein S17A/B encoding cDNA represented in SEQIDNO:40, the 40S ribosomal protein S31 encoding cDNA represented in SEQIDNO:42, the 40S ribosomal protein S10 encoding cDNA represented in SEQIDNO:44, the 60S ribosomal protein L37A encoding cDNA represented in SEQIDNO:46, the 60S ribosomal protein L34 encoding cDNA represented in SEQIDNO:48, or the 40S ribosomal protein S16 encoding cDNA represented in SEQIDNO:50.

Promoters from these or other highly expressed genes can be picked up by the method according to the invention using only routine skills of (a) making a cDNA library on mRNA isolated from a Phaffia strain grown under desired conditions, (b) determining (part of) the nucleotide sequence of the (partial) cDNAs obtained in step (a), (c) comparing the obtained sequence data in step (b) to known sequence data, such as that stored in electronic databases, (d) cloning putative promoter fragments of the gene located either directly upstream of the open reading frame or directly upstream of the transcription start site of the gene corresponding to the expressed cDNA, and (e) verifying whether promoter sequences have been obtained by expressing a suitable marker, such as the G418 resistance gene, or a suitable non-selectable "reporter" sequence downstream from a fragment obtained in (d), transforming the DNA into a Phaffia rhodozyma strain and determining the level of expression of the marker gene or reporter sequence of transformants. A transcriptional promoter is said to be of a highly expressed gene if it is capable of making Phaffia rhodozyma cells transformed with a DNA construct comprising the said promoter linked uptream of the G418 resistance marker resistant to G418 in concentrations exceeding 200 µg per liter culture medium, preferably at least 400, more prefereably more than 600 µg/l. Especially preferred promoters are those conferring resistance against more than 800 μg/ml G418 in the growth medium.

Optionally, the transcriptional start site may be determined of the gene corresponding to the cDNA corresponding to a highly expressed gene, prior to cloning the putative promoter sequences; this may serve to locate the transcriptional initiation site more precisely, and moreover, helps to determine the length of the 5'-non-translated leader of the gene. To determine the location of the transcription start site, reverse primer extension, or classical SI-mapping may be performed, based on the knowledge of the cDNA sequence. Thus the exact location of the transcription promoter can be determined without undue burden, and the isolation of a fragment upstream of the transcription start site and containing the promoter, from a hybridising genomic clone (for example a phage or cosmid) is routine. Cloning the putative promoter fragment in front (upstream) of the coding region of, for example the G418-resistance gene, and transforming the gene cassette to *Phaffia* in order to evaluate the level of G418 resistance, and hence the level of expression of the G418-resistance gene as a consequence of the presence of the promoter is routine.

In a manner essentially as described for the isolation of other strong promoters, above, a transcription terminator may be isolated, with the proviso, that the terminator is located downstream

from the open reading frame. The transcription stop site can be determined using procedures which are essentially the same as for the determination of the transcription start site. All these procedures are well known to those of skill in the art. A useful handbook is Nucleic Acid Hybridisation, Edited by B.D. Hames & S.J. Higgins, IRL Press Ltd., 1985; or Sambrook, sub. However, it is not critical that the transcription terminator is isolated from a highly expressed *Phaffia* gene, as long as it is from an expressed gene.

Using recombinant DNA according to the invention wherein the open reading frame codes for reduced sensitivity against G418, a transformation frequency was obtained up to 160 transformants per µg of linear DNA, at a G418 concentration in the medium of 40 µg/ml.

About 10 to 20 times as much transformed colonies were obtained with the vector according to the invention (pPR2) than with the prior art vector pGB-Ph9, disclosed in EP 0 590 707 A1 (see Table 2; in the experiment of Example 7, the improvement is even more striking).

The method according to the invention calls for conditions conducive to uptake of the recombinant DNA. Such conditions have been disclosed in EP 509 707. They include but are not limited to the preparation of protoplasts using standard procedures known to those of skill in the art, and subsequent incubation with the recombinant DNA. Alternatively, *Phaffia* cells may be incubated overnight in the presence of LiAc and recombinant DNA. Still further alternative methods involve the use of particle acceleration. According to a preferred embodiment, the conditions conducive to uptake involve electroporation of recombinant DNA into *Phaffia* cells, such as described by Faber et al., (1994, Current Genetics 25, 305-310). Especially preferred conditions comprise electroporation, wherein the recombinant DNA comprises *Phaffia* ribosomal DNA, said recombinant DNA being in the linear form, most preferably by cleaving said recombinant DNA in the said ribosomal region. Still further preferred conditions, comprise the use of recombinant DNA in amounts of between 1 and 10 µg per 10⁸ cells, more preferably about 5µg recombinant DNA is used per 2x10⁸ cells,

which are cultivated for 16 h at 21°C.

Once cells have been transformed according to the method, identification of transformed cells may take place using any suitable technique. Thus, identification may be done by hybridisation techniques, DNA amplification techniques such a polymerase chain reaction using primers based on the recombinant DNA used, and the like. A preferred method of identifying transformed cells is one which employs selection for the recombinant DNA that comprises a gene coding for reduced sensitivity against a selective agent. A useful selective agent is G418, hygromycin, phleomycin and amdS. Genes that code for reduced sensitivity against these selective agents are well known in the art. The open reading frames of these genes may be used as the heterologous downstream sequence according to the invention, allowing selective enrichment of transformed cells, prior to identification of transformed cells. Once transformed cells have been identified they may used for further manipulation, or used directly in the production of valuable compounds, preferably in large scale fermentors.

It will be clear, that a very efficient method for transforming *Phaffia* strains has been disclosed. Moreover, not only the frequency of transformation is high, the expression levels of the transforming DNA is very high as well, as is illustrated by the exceptionally high resistance against

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G418 of the transformed *Phaffia* cells when the open reading frame of the G418-resistance gene was fused to a promoter according to the invention when compared to the G418 resistance gene under control of the actin promoter in pGB-Ph9. It is concluded, therefore, that the GAPDH-promoter is a high-level transcriptional promoter that can be suitably used in conjunction with any heterologous DNA sequence, in order to reach high expression levels thereof in *Phaffia* strains.

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It will be clear that the availability of new expression tools, in the form of the recombinant DNA according to the invention, creates a wealth of possibilities for producing new and valuable biomolecules in *Phaffia*.

Preferably, the downstream sequence comprises an open reading frame coding for proteins of interest. For example genes already present in Phaffia, such as those involved in the carotenoid pathway. may be manipulated by cloning them under control of the high-level promoters according to the invention. Increased expression may change the accumulation of intermediates and/or end-products or change the pathway of B-carotene, cantaxanthin, astaxanthin and the like. The overexpression of the crtB gene from Erwinia uredovora will likely increase astaxanthin levels, as the product of this gene is involved in the rate limiting step. The expression of a protein of interest may also give rise to xanthophylls not known to be naturally produced in Phaffia, such as zeaxanthin. An open reading frame that may be suitably employed in such a method includes but is not limited to the one encoding the protein producing zeaxanthin (crtZ gene) obtained from Erwinia uredovora (Misawa et al.1990. J.Bacteriol. 172: 6704-6712). Other carotenoid synthesis genes can be obtained for example from Flavobacterium (a gram-positive bacterium), Synechococcus (a cyanobacterium) or Chlamydomonas or Dunaliella (algae). Obviously, carotenoid synthesis genes of a Phaffia strain, once the genes have been isolated and cloned, are suitably cloned into a recombinant DNA according to the invention and used to modify the carotenoid content of Phaffia strains. Examples of cloned carotenoid genes that can suitably be overexpressed in Phaffia, are those mentioned in Fig. 8. Particularly useful is crtE from Phycomyces blakesleanus, encoding Geranylgeranyl Diphosphate Synthase, and crtB, encoding phytoene synthase, as this step appears to be the rate-limiting step in carotenoid synthesis in Thermus thermophylus (Hoshino T. et al., 1994, Journal of Fermentation and Bioengineering 77, No. 4, 423-424). Especially preferred sources to isolate carotenoid biosynthetic genes or cDNAs from are the fungi Neurospora crassa, Blakeslea trispora. Other yeasts shown to possess cross-hybrising species of carotenoid biosynthetic genes are Cystofylobasidium, e.g. bisporidii and capitatum.

Carotenoid biosynthesis genes have also been identified in plants; these plant cDNAs or genes from plants may be used as well. Optionally, the codon usage of the Phaffia genes or cDNAs may be adapted to the preferred use in the host organism.

Of special interest according to the present invention, are the DNA sequences coding for four different enzymes in the carotenoid biosynthesis pathway of *Phaffia rhodozyma*, represented in the sequence listing. It will be clear to those having ordinary skill in the art, that once these DNA sequences have been made available it will be possible to bring about slight modifications to the DNA sequence without modifying the amino acid sequence. Such modifications are possible due to the degeneracy of the genetic code. Such modifications are encompassed in the present invention. However, also

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modifications in the coding sequences are envisaged that create modifications in the amino acid sequence of the enzyme. It is well known to those of skill in the art that minor modifications are perfectly permissible in terms of enzymatic acitivity. Most changes, such as delections, additions or amino acid substitutions do not affect enzymatic acitivity, at least not dramatically. Such variants as comprise one or more amino acid deletions, additions or substitutions can readily be tested using the complementation test disclosed in the specification. The skilled person is also familiar with the term "conservative amino acid substitutions", meaning substitutions of amino acids by similar amino acids residing in the same group. The skilled person is also familiar with the term "allelic variant", meaning naturally occurring variants of one particular enzyme. These conservative substitutions and allelic enzyme variants do not depart from the invention.

As stated, at the DNA level considerable variation is acceptable. Although the invention discloses four DNA sequences, as represented in SEQIDNO: 12, SEQIDNO: 14, SEQIDNO: 16, SEQIDNO: 18, SEQIDNO:20, or SEQIDNO: 22, in detail also isocoding variants of the DNA sequence represented in SEQIDNO: 12, SEQIDNO: 14, SEQIDNO: 16, SEQIDNO: 18, SEQIDNO: 20, or SEQIDNO: 22, are encompassed by the present invention. Those of skill in the art would have no difficulty in adapting the nucleic acid sequence in order to optimize codon usage in a host other than *P. rhodozyma*. Those of skill in the art would know how to isolate allelic variants of a DNA sequence as represented in SEQIDNO: 12, SEQIDNO: 14, SEQIDNO: 16, SEQIDNO: 18, SEQIDNO: 20, or SEQIDNO: 22 from related *Phaffia* strains. Such allelic variants clearly do not deviate from the present invention.

Furthermore, using the DNA sequences disclosed in the sequence listing, notably SEQIDNO: 12, SEQIDNO: 14, SEQIDNO: 16 or SEQIDNO: 18, as a probe, it will be possible to isolate corresponding genes form other strains, or other microbial species, or even more remote eukaryotic species if desired, provided that there is enough sequence homology, to detect the same using hybridisation or amplification techniques known in the art.

Typically, procedures to obtain similar DNA fragments involve the screening of bacteria or bacteriophage plaques transformed with recombinant plasmids containing DNA fragments from an organism known or expected to produce enzymes according to the invention. After in situ replication of the DNA, the DNA is released from the cells or plaques, and immobilised onto filters (generally nitrocellulose). The filters may then be screened for complementary DNA fragments using a labeled nucleic acid probe based on any of the sequences represented in the sequence listing. Dependent on whether or not the organism to be screened for is distantly or closely related, the hybridisation and washing conditions should be adapted in order to pick up true positives and reduce the amount of false positives. A typical procedure for the hybridisation of filter-immobilised DNA is described in Chapter 5, Table 3, pp. 120 and 121 in: Nucleic acid hybridisation- a practical approach, B.D. Hames & S.J. Higgins Eds., 1985, IRL Press, Oxford). Although the optimal conditions are usually determined empirically, a few useful rules of thumb can be given for closely and less closely related sequences.

In order to identify DNA fragments very closely related to the probe, the hybridisation is performed as described in Table 3 of Hames & Higgins, supra, (the essentials of which are reproduced

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below) with a final washing step at high stringency in 0.1 * SET buffer (20 times SET = 3M NaCl, 20 mM EDTA, 0.4 M Tris-HCl, pH 7.8), 0.1% SDS at 68° Celsius).

To identify sequences with limited homology to the probe the procedure to be followed is as in Table 3 of Hames & Higgins, supra, but with reduced temperature of hybridisation and washing. A final wash at 2 * SET buffer, 50°C for example should allow the identification of sequences having about 75% homology. As is well known to the person having ordinary skill in the art, the exact relationship between homology and hybridisation conditions depend on the length of the probe, the base composition (% of G + C) and the distribution of the mismatches; a random distribution has a stronger decreasing effect on T_m then a non-random or clustered pattern of mismatches.

The essentials of the procedure described in Table 3, Chapter 5 of Hames & Higgins are as follows:

(1) prehybridisation of the filters in the absence of probe, (2) hybridisation at a temperature between 50 and 68°C in between 0.1 and 4 * SET buffer (depending on the stringency), 10 * Denhardt's solution (100 * Denhardt's solution contains 2% bovine serum albumin, 2% Ficoli, 2% polyvinylpyrrolidone), 0.1% SDS, 0.1% sodiumpyrophosphate, 50 μg/ml salmon sperm DNA (from a stock obtainable by dissolving 1 mg/ml of salmon sperm DNA, sonicated to a length of 200 to 500 bp, allowed to stand in a water bath for 20 min., and diluted with water to a final concentration of 1 mg/ml); hybridisation time is not too critical and may be anywhere between 1 and 24 hours, preferably about 16 hours (o/n); the probe is typically labeled by nick-translation using ³²P as radioactive label to a specific activity of between 5 * 10⁷ and 5 * 10⁸ c.p.m./μg; (3) (repeated) washing of the filter with 3 * SET, 0.1% SDS, 0.1% sodiumpyrophosphate at 68°C at a temperature between 50°C and 68°C (dependent on the stringency desired), repeated washing while lowering the SET concentration to 0.1%., wash once for 20 min. in 4 * SET at room temperature, drying filters on 3MM paper, exposure of filters to X-ray film in a cassette at -70°C for between 1 hour and 96 hours, and developing the film.

Generally, volumina of prehybridisation and hybridisation mixes should be kept at a minimum. All "wet" steps may be carried out in little sealed bags in a pre-heated water bath.

The above procedure serves to define the DNA fragments said to hybridise according to the invention. Obviously, numerous modifications may be made to the procedure to identify and isolate DNA fragments according to the invention. It is to be understood, that the DNA fragments so obtained fall under the terms of the claims whenever they can be detected following the above procedure, irrespective of whether they have actually been identified and/or isolated using this procedure.

Numerous protocols, which can suitably be used to identify and isolate DNA fragments according to the invention, have been described in the literature and in handbooks, including the quoted Hames & Higgins, supra).

With the advent of new DNA amplification techniques, such as direct or inverted PCR, it is also possible to clone DNA fragments in vitro once sequences of the coding region are known.

Also encompassed by the claims is a DNA sequence capable, when bound to nitrocellulose filter and after incubation under hybridising conditions and subsequent washing, of specifically hybridising to a radio-labelled DNA fragment having the sequence represented in SEQIDNO: 12,

SEQIDNO: 14, SEQIDNO: 16 or SEQIDNO: 18, as detectable by autoradiography of the filter after incubation and washing, wherein said incubation under hybridising conditions and subsequent washing is performed by incubating the filter-bound DNA at a temperature of at least 50°C, preferably at least 55°C, more preferably at least 60°C in the presence of a solution of the said radio-labeled DNA in 0.3 M NaCl, 40 mM Tris-HCl, 2 mM EDTA, 0.1% SDS, pH 7.8 for at least one hour, whereafter the filter is washed at least twice for about 20 minutes in 0.3 M NaCl, 40 mM Tris-HCl, 2 mM EDTA, 0.1% SDS, pH 7.8, at a temperature of 50°C, preferably at least 55°C, more preferably at least 60°C, prior to autoradiography.

The heterologous DNA sequence according to the invention may comprise any open reading frame coding for valuable proteins or their precursors, like pharmaceutical proteins such as human serum albumin, IL-3, insulin, factor VIII, tPA, EPO, α-interferon, and the like, detergent enzymes, such as proteases and lipases and the like, cell wall degrading enzymes, such as xylanases, pectinases, cellulases, glucanases, polygalacturonases, and the like, and other enzymes which may be useful as additives for food or feed (e.g. chymosin, phytases, phospholipases, and the like). Such genes may be expressed for the purpose of recovering the protein in question prior to subsequent use, but sometimes this may not be necessary as the protein may be added to a product or process in an unpurified form, for example as a culture filtrate or encapsulated inside the *Phaffia* cells.

The yeast cells containing the carotenoids can be used as such or in dried form as additives to animal feed. Furthermore, the yeasts can be mixed with other compounds such as proteins, carbohydrates or oils.

Valuable substances, such as proteins or pigments produced by virtue of the recombinant DNA of the invention may be extracted. Carotenoids can also be isolated for example as described by Johnson et al. (Appl. Environm. Microbiol. 35: 1155-1159 (1978)).

Purified carotenoids can be used as colorants in food and/or feed. It is also possible to apply the carotenoids in cosmetics or in pharmaceutical compositions.

The heterologous downstream sequence may also comprise an open reading frame coding for reduced sensitivity against a selective agent. The open reading frame coding for an enzyme giving G418 resistance was used satisfactorily in the method according to the invention, but the invention is not limited to this selection marker. Other useful selection markers, such as the phleomycin resistance gene may be used, as disclosed in EP 590 707. Each of these genes is advantageously expressed under the control of a strong promoter according to the invention, such as the GAPDH-promoter.

The invention is now being illustrated in greater detail by the following non-limitative examples.

Experimental

Strains: E. coli DH5a: supE44lacU169 (80lacZM15) hsdR17 recA1 endA1 gyrA96 thi-1 relA1

E. coli LE392: supE44 supF58 hsdR514 galK2 galT22 metB1 trpR55 lacY1

P. rhodozyma CBS6938

Plasmids:

pUC19 (Gibco BRL)

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pTZ19R

PUC-G418

pGB-Ph9 (Gist-brocades)

pMT6 (1987, Breter H.-J., Gene 53, 181-190))

Media: LB: 10 g/l bacto tryptone, 5 g/l yeast extract, 10 g/l NaCl. Plates; +20 g/l bacto agar. When appropriate 50 μg/ml ampicillin.

YePD: 10 g/l yeast extract, 20 g/l bacto peptone, 20 g/l glucose. Plates; +20 g/l bacto agar. When appropriate 50 µg/ml Geneticin (G418).

Methods: All molecular cloning techniques were essentially carried out as described by Sambrook et al. in Molecular Cloning: a Laboratory Manual, 2nd Edition (1989; Cold Spring Harbor Laboratory Press).

Enzyme incubations were performed following instructions described by the manufacturer. These incubations include restriction enzyme digestion, dephosphorylation and ligation (Gibco BRL).

Isolation of chromosomal DNA from *Phaffia rhodozyma* as described in example 3 of patent Gist-brocades; EP 0 590 707 A1. Chromosomal DNA from *K. lactis* and *S.cerevisiae* was isolated as described by Cryer et al. (Methods in Cell Biology 12: 39, Prescott D.M. (ed.) Academic Press, New York)

Isolation of large (> 0.5-kb) DNA fragments from agarose was performed using the Geneclean II Kit whereas small (< 0.5-kb) and DNA fragments or fragments from PCR mixtures were isolated using WizardTM DNA Clean-Up System (Promega).

Transformation of *E. coli* was performed according to the CaCl₂ method described by Sambrook *et al.* Packaging of cosmid ligations and transfection to *E. coli* LE392 was carried out using the Packagene Lambda DNA Packaging System (Promega), following the Promega protocols.

Isolation of plasmid DNA from E. coli was performed using the QIAGEN (Westburg B.V. NL).

Transformation of *Phaffia* CBS6938 was done according to the method for *H. polymorpha* described by Faber *et al.*, *supra*;

- Inoculate 30 ml of YePD with 1 CBS6938 colony
- Grow 1-2 days at 21°C, 300 rpm (pre-culture)
- Inoculate 200 ml of YePD with pre-culture to OD₆₀₀ = between 0 and 1 (if above 1 dilute with water)
- Grown o/n at 21°C, 300 rpm until OD₈₀₀ = 1.2 (dilute before measuring)
 - Centrifuge at 5 min. 8000 rpm, room temperature. Remove supernatant thoroughly
 - Resuspend pellet in 25 ml 50 mM KPi pH 7.0, 25 mM DTT (freshly made)

Transfer suspension to a fresh sterile 30 ml centrifuge tube and incubate for 15 min. at room temperature

- Centrifuge 5 min. at 8000 rpm 4°C, remove supernatant thoroughly
- Resuspend pellet in 25 ml of ice cold STM (270 mM sucrose, 10 mM Tris pH 7.5, 1 mM MgCl₂)
 - Centrifuge 5 min. at 8000 rpm, 4°C
 - Repeat washing step
 - Resuspend cells in 0.5 ml of ice cold STM (3*109 cells/ml). Keep on ice!

- Transfer 60 µl of cell suspension to pre-cooled Eppendorf tubes containing 5 µg transforming DNA (use precooled tips!), Keep on ice
- -Transfer Cell/DNA mix to precooled electroporation cuvettes (top to bottom)
- Pulse: 1.5 kV, 400 Ω, 25 μF
- Immediately add 0.5 ml of ice cold YePD. Transfer back to ep using a sterile Pasteur pipette
 - Incubate 2.5 hrs at 21°C
 - Plate 100 μl onto YePD-plates containing 40 μg/ml G418
 - Incubate at 21°C until colonies appear.

Pulsed Field Electrophoresis was performed using a GENE Navigator + accessories (Pharmacia). Conditions: 0.15 * TBE, 450 V, pulse time 0.5 s, 1.2% agarose, run time 2 h.

Polymerase Chain Reaction (PCR) experiments were performed in mixtures having the following composition:

- 5 ng of plasmid DNA or 1 µg chromosomal DNA
- 0.5 μ g of oligo nucleotides (5 μ g degenerated oligo's in combination with chromosomal DNA)
- 10 nm of each dNTP
- 2.5 µm KCl
- 0.5 μm Tris pH 8.0
- 0.1 μm MgCl2
- 0.5 μg gelatin
- 1.3 U Taq polymerase (5 U in combination with chromosomal DNA)

 H_2O was added to a total volume of 50 μ l

Reactions were carried out in an automated thermal cycler (Perkin-Elmer).

Conditions: 5 min. 95°C, followed by 25 repeated cycli; 2' 94°C, 2' 45°C3' 72°C

25 Ending; 10 min. 72°C.

Fusion PCR reactions were performed as described above, except that 2 DNA fragments with compatible ends were added as a template in equimolar amounts.

Oligo nucleotide sequences were as follows:

3005: CGGGATCCAA(A/G)CTNACNGGNATGGC (SEQIDNO: 1);

3006: CGGGATCC(A/G)TAICC(C/A/G)(C/T)A(T/C)TC(A/G)TT(A/G)TC(A/G)TACCA (SEQIDNO: 2);

4206: GCGTGACTTCTGGCCAGCCACGATAGC (SEQIDNO: 3);

5126: TTCAATCCACATGATGGTAAGAGTGTTAGAGA (SEQIDNO: 4);

5127: CTTACCATCATGTGGATTGAACAAGATGGAT (SEQIDNO: 5);

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5177: CCCAAGCTTCTCGAGGTACCTGGTGGGTGCATGTATGTAC (SEQIDNO: 6);

5137: CCAAGGCCTAAAACGGATCCCTCCAAACCC (SEQIDNO: 7);

5138: GCCAAGCTTCTCGAGCTTGATCAGATAAAGATAGAGAT (SEQIDNO: 8);

Example 1 G-418 resistance of Phaffia transformant G418-1

To determine the expression of the G418 resistance gene in pGB-Ph9, transformant G418-1 (EP 0 590 707 A1) was exposed to increasing concentrations of G418.

Two dilutions of a G418-1 culture were plated onto YepD agar containing 0-1000 µg/ml G418 (Table 1).

	[G418] μg/ml	Phaffia G418-1 Dil.=10⁴(OD ₆₀₀ =7)	Phaffia G418-1 Dil.=10 ⁻⁵ (OD ₆₀₀ =7)	Phaffia (CBS6938) Dil.=0(OD ₆₀₀ =5)
15	0	>300	74	>300
	200	>300	70	0
	300	>300	61	0
	400	212	13	0
	500	10	2	0
20	600	0	0	0
	700	0	0	0
	800	0 .	0	0
	900	0	0	0
	1000	0	0	0

Table 1. Survival of Phaffia transformant G418-1 on YepD agar medium containing increasing concentrations of G418.

At a concentration of 600 μg/ml G418 less than 1% of the plated cells survived. It can be concluded, that despite multicopy integration of pGB-Ph9, G418-1 shows a rather weak resistance to G418 (Scorer et al., 1994, Bio/Technology 12, p. 181 et seq., Jimenez and Davies, 1980, Nature 187 p. 869 et seq.), most probably due to a weak action of the Phaffia actin promoter in the plasmid. The results that the Phaffia actin promoter works poorly, prompted us to isolate promoter sequences of Phaffia with strong promoter activity.

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Example 2

Synthesis of specific probes of glycolytic genes from Phaffia rhodozvma by PCR

The polymerase chain reaction (PCR) technique was used in an attempt to synthesize a homologous probe of the genes encoding glyceraldehyde-3-phosphate dehydrogenase (GAPDH), phosphoglycerate kinase (PGK) and the triose phosphate isomerase (TPI) of *Phaffia rhodozyma*.

A set of degenerated oligonucleotides was designed based on the conserved regions in the GAPDH-gene (Michels et al., 1986. EMBO J. 5: 1049-1056), PGK-gene (Osinga et al., 1985. EMBO J. 4: 3811-3817) and the TPI-gene (Swinkels et al., 1986. EMBO J. 5: 1291-1298).

All possible oligo combinations were used to synthesize a PCR-fragment with chromosomal DNA of *Phaffia rhodozyma* (strain CBS6938) as template. Chromosomal DNA of *Saccharomyces cerevisiae* and *Kluvveromyces lactis* as template was used to monitor the specificity of the amplification. The PCR was performed as described above, the PCR conditions were 1' 95 °C, 2' annealing temperature (T_a), in 5' from annealing temperature to 72 °C, 2' 72 °C, for 5 cycli followed by 1' 95 °C, 2' 55 °C and 2' 72 °C for 25 cycli and another elongation step for 10' 72 °C. Three different T_a were used 40 °C, 45 °C and 50 °C.

Under these conditions, only one primer combination produced a fragment of the expected size on chromosomal DNA of *Phaffia* as template. Using the oligo combination no: 3005 and 3006 and a T_a of 45 °C a 0.3-kb fragment was found. Specifically, the GAPDH oligonucleotides correspond with amino acids 241-246 and 331-338 of the published *S. cerevisiae* sequence. (It was concluded that to isolate the promoters corresponding to the PGK- and TPI-genes from *Phaffia*, either further optimization of the PCR-conditions is required, or homologous primers should be used. Another alternative method for isolating high level promoters is disclosed in the detailed description, *supra*.

The amplified fragment was purified from the PCR reaction and was digested with BamHI and ligated into the dephosphorylated BamHI site of pTZ19R. The ligation mixture was transformed to competent E. coli DH5α cells prepared by the CaCl₂-method and the cell were plated on LB-plates with 50 μg/ml Amp and 0.1 mM IPTG/50 μg/ml X-gal. Plasmid DNA was isolated from the white colonies. The pTZ19R clone with the right insert, called pPRGDH1, was subsequently used for sequence analysis of the insert.

The cloned sequence encoded for the carboxy terminal fragment of GAPDH of *Phaffia* as shown by comparison with the GAPDH-gene sequence of S. cerevisiae (Holland and Holland, 1979. J. of Biol. Chem. 254: 9839-9845).

Example 3

Isolation of the GAPDH-gene of Phaffia

To obtain the complete GAPDH-gene including expression signals the 0.3-kb <u>Bam</u>HI fragment of pPRGDH1 was used to screen a cosmid library of *Phaffia*.

Preparation of the vector for cosmid cloning.

Vector preparation was simplified, because of the presence of a double cos-site in pMT6. PMT6 was digested to completion with blunt end cutter *Pvull* to release the cos-sites. Digestion efficiency was checked by transformation to *E. coli* DH5α and found to be >99%.

The Pvull digested pMT6 was purified by phenol:chloroform extraction and ethanol precipitation and finally solved in 30 μ l TE at a concentration of 2 μ g/ μ l.

The vector was subsequently digested with cloning enzyme BamHI and the vector arms were purified as described above ("Experimental").

Preparation of target DNA

Isolation of genomic DNA of *Phaffia* strain CBS6938 was performed as described in the part named "Experimental". The cosmid pMT6 containing inserts of 25-38-kb are most efficiently packaged. Therefore genomic DNA was subjected to partial digestion with the restriction enzyme *Sau*3A. Target DNA was incubated with different amounts of enzyme. Immediately after digestion the reactions were stopped by the extraction of DNA from the restriction mixture with phenol-chloroform. The DNA was precipitated by using the ethanol method and the pelleted DNA after centrifugation was dissolved in a small volume of TE. Contour clamped homogeneous electric field (CHEF) electrophoresis was used to estimate the concentration and size of the fragments (Dawkins, 1989, J. of Chromatography 492, pp. 615-639).

20 Construction of genomic cosmid library.

Ligation of approximately 0.5 μ g of vector arm DNA and 0.5 μ g of target DNA was performed in a total volume of 10 μ l in the presence of 5 mM ATP (to prevent blunt end ligation).

Packaging in phage heads and transfection to E. coli LE 392 as described in Experimental.

The primary library consisted of 7582 transfectants with an average insert of 28-kb as determined by restriction analysis. The library represents 3.5 times the genome with a probability of the presence of all genes in the library of 0.97 as calculated according to Sambrook (supra). For library amplification the transfectants were pooled by resuspending in 8 ml LB-broth. Additional 4.8 ml glycerol was added. The transfectants mixture was divided into 16 samples of 800 µl each and stored at -80 °C. This amplified library consisted of 2.9*10° transfectants.

Screening of the cosmid library.

A 100 µl sample was taken from this library and further diluted (106) in LB-broth and 200 µl was plated onto 10 LB-plates containing ampicillin. The plates were incubated overnight at 37 °C. Each plate contained 300-400 colonies and filters were prepared. These filters were screened with the GAPDH-probe using hybridization and washing conditions as described above ("Experimental"). After 16 hours exposure, 3 strong hybridization signals were found on the autoradiogram.

Cosmid DNA isolated from these positive colonies was called pPRGDHcos1, pPRGDHcos2 and pPRGDHcos3.

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Chromosomal DNA isolated from *Phaffia rhodozyma* strain CBS 6938 and cosmid pPRGDHcos1 was digested with several restriction enzymes. The DNA fragments were separated, blotted and hybridized as described before. The autoradiograph was exposed for different time periods at -80°C. The film showed DNA fragments of different length digested by different restriction enzymes which hybridize with the GAPDH-probe (Fig. 1).

Furthermore, from Southern analysis of the genomic DNA of *Phaffia* using the GAPDH fragment as probe, it was concluded that the GAPDH-encoding gene is present as a single copy gene in *Phaffia rhodozyma*, whereas in *Saccaromyces cerevisiae* GAPDH is encoded by three closely related but unlinked genes (Boucherie *et al.*, 1995. FEMS Microb. Letters <u>135</u>:127-134).

Hybridizing fragments of pPRGDHcos1 for which a fragment of the same length in the chromosomal DNA digested with the same enzyme was found, were isolated from an agarose gel. The fragments were ligated into the corresponding sites in pUC19. The ligation mixtures were transformed to competent *E. coli* cells. The plasmids with a 3.3-kb *Sal*1 insert and a 5.5-kb *Eco*Rl insert were called pPRGDH3 and pPRGDH6, respectively. The restriction map of pPRGDH3 and pPRGDH6 is shown in Figure 2. Analysis of the sequence data of the insert in pPRGDH1 showed us that there was a *Hindlll* site at the C-terminal part of the GAPDH-gene. From this data it was suggested that the insert in pPRGDH6 should contain the complete coding sequence of GAPDH including promoter and terminator sequences.

Example 4

Characterization of the GAPDH-gene

In order to carry out sequence analysis without the need to synthesize a number of specific sequence primers a number of deletion constructs of plasmids pPRGDH3 and pPRGDH6 were made using convenient restriction sites in or near the putative coding region of GAPDH gene.

The plasmids were digested and after incubation a sample of the restriction mixture was analyzed by gel electrophoresis to monitor complete digestion. After extraction with phenol-chloroform the DNA was precipitated by ethanol. After incubation at -20 °C for 30' the DNA is pelleted by centrifugation, dried and dissolved in a large volume (0.1 ng/μl) of TE. After ligation the mixtures were transformed to E. coli. Plasmid DNA isolated from these transformants was analyzed by restriction analysis to reveal the right constructs. In this way the deletion constructs pPRGDH3δHIII, pPRGDH6δBamHI, pPRGDH6δSstl and pPRGDH6δSall (Fig. 1).

In addition to this, the 0.6-kb and 0.8-kb Sstl fragments derived from pPRGDH6 were subcloned in the corresponding site of pUC19.

Sequence analysis was carried out using pUC/M13 forward and reverse primers (Promega). The sequencing stategy is shown in fig. 2 (see arrows).

On the basis of homology with the GAPDH-gene sequence of *S. cerevisiae* (Holland and Holland, 1979. J. of Biol. Chem. <u>254</u>: 9839-9845) and *K. lactis* (Shuster, 1990. Nucl. Acids Res. <u>18</u>, 4271) and the known splice site concensus J.L. Woolford. 1989. Yeast <u>5</u>: 439-457), the introns and the possible ATG start were postulated.

The GAPDH gene has 6 introns (Fig. 1) and encodes a polypeptide of 339 amino acids. This was completely unexpected considering the genomic organisation of the GAPDH genes of *K. lactis* and *S. cerevisiae* which have no introns and both consist of 332 amino acids. The homology on the amino acid level between the GAPDH gene of *Phaffia* and *K. lactis* and *S. cerevisiae* is 63% and 61%, respectively.

Most of the introns in the GAPDH gene are situated at the 5' part of the gene. Except intron III all introns contain a conserved branch-site sequence 5'-CTPuAPy-3' found for S. cerevisiae and S. pombe.

By computer analysis of the upstream sequence using PC-gene 2 putative eukaryotic promoter elements, TATA-box (position 249-263 in SEQIDNO: 11) and a number of putative Cap signal (between position 287 and 302 in SEQIDNO: 11) were identified.

Example 5

Cloning of the GAPDH promoter fused to G418 in pUCG418.

In order to construct a transcription fusion between the GAPDH promoter and the gene encoding G418 resistence the fusion PCR technique was used.

Using plasmid pPRGDH6 the GAPDH promoter could be amplified by standard PCR protocols ("Experimental").

In the PCR mix pPRGDH6 and oligo's No. 5177 and 5126 (Sequences in "Experimental") were used. A 416 bp DNA fragment was generated containing the entire GAPDH promoter sequence. In addition this fragment also contains a *HindIII*, *XhoI* and a *KpnI* restriction site at it's 5'end and 12 nt overlap with the 5' end of the gene encoding G418 resistance.

The 217 bp portion of the 5'end of the G418 coding sequence was also amplified by PCR using pUC-G418 and oligo's 4206 and 5127. A 226 bp DNA fragment was obtained containing the 217 bp 5'end of G418 and having a 9 nucleotides overlap with the 3'end of the earlier generated GAPDH promoter fragment. It also contained a *Mscl* site at it's 3end.

The PCR fragments were purified from the PCR mixture using the WIZARD Kit.

Approximately 1 µg of the GAPDH promoter fragment and 1 µg of the G418 PCR fragment were used together with oligo's 5177 and 4206 in a fusion PCR experiment (Experimental). A 621 bp DNA fragment was generated, containing the GAPDH promoter directly fused to the 5' portion of G418. After purification the DNA fragment was digested with Msc1 and Kpn1. The 3.4 Kb Msc1-Kpn1 fragment of pUC-G418, containing pUC sequences and the 3' portion of G418, was used as a vector.

The ligation mixture was transformed to competent E. coli DH5 α cells. Transformant colonies containing the fusion PCR DNA inserted were identified by digestion with different restriction enzymes.

Thus, plasmid pPR1 was obtained, containing the GAPDH promoter directly fused to the G418 marker gene. Three pPR1 vectors isolated from independent transformants were used in further cloning experiments.

To target the plasmid, after transformation, to a specific integration site a 3.0-kb Sst1 fragment containing a part of the ribosomal DNA of *Phaffia* was cloned in pPR1. The ribosomal DNA fragment was isolated from an agarose gel after digestion with Sst1 of plasmid pGB-Ph11 (EP 590 707 A1). This

fragment was ligated in the dephosphorylated Sstl site of pPR1. The ligation mixture was transformed to competent *E. coli* cells. Plasmid DNA was isolated and using restriction analysis it was shown that several colonies contain the expected plasmid pPR2. The complete cloning strategy is shown in Fig. 3.

Example 6

Transformation of Phaffia with pPR2.

Transformation of *Phaffia* strain 6938 was performed using an electroporation procedure as previously described by Faber et al. (1994, Curr. Genet. 1994: 25,305-310) with the following modifications:

- Electropulsing was performed using the Bio-rad Gene Pulser with Pulse Controller and with Bio-rad 2mm cuvettes.
 - Phaffia was cultivated for 16 h at 21 °C.
 - Per transformation $2x10^8$ cells were used together with 5 µg of linearized vector. Linearization was done in the rDNA sequence using Clal to enable integration at the rDNA locus in the Phaffia genome. Following the electric pulse (7.5kV/cm, 400 Ω and 25 µF) 0.5 ml YePD medium was added to the cell/DNA mixture. The mixture was incubated for 2.5 h at 21 °C and subsequently spread on 5 selective YEDP agar plates containing 40 µg/ml G418.

As shown in Table 2 we were able to generate transformants with 115 transformants per µg DNA; the average transformation frequency was 50 transformants/µg pPR2 as judged over a number of experiments. Transformation of the closed circular form of pPR2 did not result in transformation suggesting that there is no autonomously replicating sequence present within the vector sequences. Using pPR2 a 10 to 50-fold increase in transformation frequency was found compared to a previous constructed transformation vector for *Phaffia*, called pGB-Ph9. In this latter vector a translation fusion was made between the 5' part of the actin gene of *Phaffia* and G418.

In order to analyze the level of resistance of transformants the mixture or DNA/cells was plated onto selective plates containing different amounts of G418. Although the total number of transformants decreases with the increasing amounts of G418, we were still able to obtain a considerable number of transformants (table 3).

In another experiment 30 transformants obtained under standard selection conditions (40 μ g/ml) were transferred to plates containing 50, 200 or 1000 μ g/ml. After incubation of the plates at 21 °C for 4-5 days, 23 transformants out of 30 tested were able to grow on plates containing 200 μ g/ml G418. One transformant was able to grow on plates containing upto and above 1000 μ g/ml G418.

	Table 2.	Transforma	tion frequency of pGB-Ph9 and pPR2
35		Exp. I	Exp.2
		69	8
	pGB-Ph9x <i>Bgl</i> II	46	7
40	pPR2 ccc	n.d	n.d
	pPR2(A)xClal	714	56
	(B)	639	124

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(C) 443 153

Total number of transformants (> 1 mm) in different transformation experiments after 4-5 days incubation.

Table 3. Comparison of G418 sensitivity as a result of two different G418-resistance genes in pGB-Ph9 and pPR2

10	concentration G418 (µg/ml)	Number of transformants	
		pPR2x <i>Cla</i> l	pGB-Ph9xBg/II (=pYac4)
15	40	480	2
	50	346	-
	60	155	•
	70	61	-
	80	141	<u>.</u> *
20	90	72	-
	100	64	-

Analysis of pPR2 transformants.

To analyse the integration event and the number of integrated vector copies total genomic DNA from six independent transformants was isolated. Therefore these transformants were cultivated under selective conditions, i.e. YePD + 50 µg/ml G418. Chromosomal DNA was digested with Clal. The DNA fragments were separated by gel electrophoresis and transferred to nitrocellulose and the Southern blot was probed with Phaffia DNA.

Besides the rDNA band of 9.1 kb an additional band of 7.1 kb of similar fluorescing intensity was observed in the transformants. This band corresponds to the linearised form of pPR2. From the intensity of these bands it was concluded that the copy number was about 100 - 140 copies of pPR2. These results are similar to those observed for pGB-Ph9, ruling out that the improved G418-resistance is due to differences in copy number of integrated vectors alone. It is not known whether the multiple copy event is caused by multiple copy integration of pPR2 or by the amplification of a single copy in the rDNA or a combination of both events.

Example 7

Construction of pPR2T by cloning the GAPDH-terminator into pPR2

Eukaryotic mRNAs contain modified terminal sequences, specifically the 3' terminal poly(A). As the prokaryotic gene encoding G418 resistance lacks eukaryotic termination signals, which might effect proper transcription termination and mRNA stability (1994, Raue, H.A., TIBTECH 12: 444-449), a part of the 3' non-coding sequence of GAPDH was introduced.

To that end, a 307 bp fragment, consisting of 281 bp of the 3' non-coding region of GAPDH and other additional cloning sequences, was amplified by PCR using the oligo's 5137 and 5138 ("Experimental").

The upstream oligo 5137 consists of the last 14 nucleotides of the coding and 17 nucleotides of the 3' non-coding region of GAPDH. By base substitutions of the 5th (T --> A) and 8th (T --> C) nucleotide

of the non-coding sequence a BamHI restriction site was introduced. In addition this fragment contains a XhoI and a HindIII restriction site at its 3' end.

The PCR fragment was purified from the PCR mixture using the WIZARD Purification Kit and digested with *Bam*HI and *Hind*III. A 288 bp fragment was isolated and cloned into the corresponding sites of the previously constructed *Phaffia* transformation vector pPR2, yielding pPR2T.

Upon transformation of *Phaffia*, using G418 as selective agent, the transformation frequencies (number of transformants per µg of DNA) obtained with the improved construct pPR2T was approximately 5 to 10 times higher than the transformation frequency of pPR2 (i.e. without a *Phaffia* homologous transcription termination signal). The results of a typical experiment are given in Table 4.

Table 4 Transformation frequency at 50 μg/ml G418 for pGB-Ph9, pPR2 and pPR2T

Vector	transformants	transformants/µg DNA
pGB-Ph9 (ccc)	-	-
pGB-Ph9 (x <i>Bgl</i> II)	60	1
pPR2 (ccc)	1	-
pPR2 (xClai)	3000 - 9600	50 - 160
pPR2T (ccc)	- -	-
pPR2T (x <i>Cla</i> I)	45600	760
pPR2T (x.SfiI)	1080	18

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Phaffia cells transformed with pPR2T were tested for their ability to grow on high levels of G418. The level of G418 on which growth is still possible was taken as a measure of the expression level of the G418 resistance gene in transformants, as a result of the presence of the Phaffia promoter, and/or terminator. Preliminary results indicate that the number of transformants able to grow on high levels of G418 are significantly higher than without terminator.

In summary

From the above results, it was concluded, that the presence of the GAPDH-promoter (pPR2) resulted in a considerable increase of the transformation frequency (from 1 to at least 50 per µg of DNA) when compared to the vector containing the actin-promoter (pGB-Ph9). These results are in line with the results obtained with the G418 sensitivity test (Table 3 and 4) which indicate superior expression levels under the control of the GAPDH promoter. The possibility that the difference in transformation frequency could be due solely to the difference in linearising the vectors, (Bgl11, Cla1 and Sfi1 all cut inside the ribosomal DNA locus, but at different positions), was ruled out by comparison of pPR2(xSfi1) with pGB-Ph9(xSfi1). The difference in transformation frequency between the two pPR2 and pGB-Ph9, linearised with Sfi1 is still considerable. However, it is concluded that the choice of the linearisation site does have effect on the transformation frequency; linearisation with Cla1 is preferred.

The improvements obtained by using a high-level promoter, such as GAPDH, are irrespective of whether a homologous terminator is used (pPR2 (without homologous terminator) performs far better than pGB-Ph9, both in G418 sensitivity tests, as well as in terms of transformation frequency).

The presence of a homologous terminator results in both higher transformation frequencies and higher expression levels; this result is concluded to be independent of the promoter used. Preliminary results indicate that considerable improvements are obtained when the pGB-Ph9 construct is completed with a transcription terminator, such as the GAPDH-terminator used in pPR2T.

The following Examples illustrate the isolation of DNA encoding enzymes involved in the carotenoid biosynthesis pathway of *Phaffia rhodozyma*. These DNA sequences can suitably be used for a variety of purposes; for example to detect and isolate DNA sequences encoding similar enzymes in other organisms, such as yeast by routine hybridisation procedures, to isolate the transcription promoters and/or terminators, which can be used to construct expression vectors for both heterologous as well as homologous downstream sequences to be expressed. The DNA sequences encoding carotenoid biosynthesis genes can suitably be used to study the over-expression, either under the control of their own promoters or heterologous promoters, such as the glycolytic pathway promoters illustrated above. For example, transformation of *Phaffia rhodozyma* with carotenoid encoding DNA sequences according to the invention effectively results in amplification of the gene with respect to the wild-type situation, and as a consequence thereof to overexpression of the encoded enzyme.

Hence, the effect of over-expression of one or more genes encoding carotenoid biuosynthesis genes can thus be studied. It is envisaged that mutant Phaffia strains can be obtained producing higher amounts of valuable carotenoids, such as \(\theta\)-carotene, cantaxanthin, zeaxanthin and/or astaxanthin. Similarly, the DNA sequences encoding enzymes involved in the carotenoid biosynthesis pathway can be introduced into other hosts, such as bacteria, for example \(E\) coli, yeasts, for example species of \(Saccharomyces\), \(Kluyveromyces\), \(Rhodosporidium\), \(Candida\), \(Yarrowia\), \(Phycomyces\), \(Hansenula\), \(Picchia\), \(\text{fungi}\), \(\text{such as a carrot}\), tomato, and the like. The procedures of transformation and expression requirements are well known to persons skilled in these arts.

Strains: E. coli XL-Blue-MRF'Δ(mcrA)183Δ(mcrCB-hsdSMR-mrr) 173 endA! supE44 thi-l recAl gyrA96 relAl lac[F' proAB laq^qZΔM15 Tn10 (Tet')]

ExAssistTM interference-resistant helper phage (Stategene^R)

P. rhodozyma CBS6938 or

P. rhodozyma asta 1043-3

Plasmids used for cloning:

pUC19 Apr (Gibco BRL)

Uni-ZAP™ XR vector (lambda ZAP^R !I vector digested with *EcoRI-XhoI*, CIAP treated;Strategene^R)

Media: LB: 10 g/l bacto tryptone, 5 g/l yeast extract, 10 g/l NaCl. Plates; +20 g/l bacto agar.

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When appropriate 50-100 μ g/ml ampicillin (Ap), 30 μ g/ml chloramphenicol (Cm) and 1 mM isopropyl-1-thio- β -D-galactopyranoside (IPTG) was added.

YePD: 10 g/l yeast extract, 20 g/l bacto peptone, 20 g/l glucose. Plates; +20 g/l bacto agar.

All molecular cloning techniques were essentially carried out as described by Sambrook et al. in Molecular Cloning: a Laboratory Manual, 2nd Edition (1989; Cold Spring Harbor Laboratory Press). Transformation of E. coli was performed according to the CaCl₂ method described by Sambrook et al.

Enzyme incubations were performed following instructions described by the manufacturer. These incubations include restriction enzyme digestion, dephosphorylation and ligation (Gibco BRL). Isolation of plasmid DNA from <u>E. coli</u> was performed using the QIAGEN (Westburg B.V. NL).

For sequence analysis deletions constructs and oligonucleotides were made to sequence the complete sequence using a *Taq* DYE Primer Cycle Sequencing kit (Applied Biosystems).

Example 8

Description of plasmids

Plasmids (pACCAR25ΔcrtE, pACCAR25ΔcrtB, pACCRT-EIB, pACCAR16ΔcrtX and pACCAR25ΔcrtX), which contain different combinations of genes involved in the biosynthesis of carotenoid in *Erwinia uredovora* were gifts from Prof. Misawa; Kirin Brewery co.,LTD.; Japan). The biosynthetic route of carotenoid synthesis in *Erwinia uredovora* is shown in fig 8.

In addition a derivative of pACCAR25\(\Delta\colon\) crtX, designated pACCAR25\(\Delta\colon\) crtX\(\Delta\colon\) crtI, was made in our laboratory. By the introduction of a frameshift in the BamHI restriction site the crtI gene was inactivated. E. coli strains harboring this plasmid acumulate phytoene which can be monitored by the red phenotype of the colony.

All plasmids are derivatives of plasmid pACYC184 (Rose RE; Nucl. Acids Res. 16 (1988) 355), which contains a marker conferring chloramphenicol-resistance. Furthermore these plasmids and derivatives thereof contain a replication origin that is compatible to vectors such as pUC and pBluescript. Each plasmid contains a set of carotenoid biosynthetic genes of *Erwinia uredovora* mediating the formation of different carotenoid in *E. coli*. The complete list of plasmid used in this study is shown in Table 5.

Table 5: Summary of carotenoid producing *E.coli* strains used in this study.

PLASMID:	GENOTYPE:	CAROTENOID ACCUMULATED:	COLOR PHENOTYPE:
pACCAR25∆cπE	crtB; crtI; crtY; crtX; crtZ	farnesyl pyrophosphate/iso- pentenyl pyrophosphate	white
pACCAR25∆crtB	crtE; crtI; crtY; crtX; crtZ	geranylgeranyl pyrophosphate	white
pACCAR25AcrtX Acrtl	crtE; crtB; crtY; crtZ	phytoene	white

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pACCRT-EIB	crtE; crtB; crtI	lycopene	red
pACCAR16ΔcπX	crtE; crtB; crtI crtY	β-carotene	yellow
pACCAR25ΔετίΧ	crtE; crtB; crtI; crtY; crtZ	zeaxanthin	yellow/ orange

Genes encoding: crtE, geranylgeranyl pyrophosphate synthase; crtB, Phytoene synthase; crtI, phytoene desaturase; crtI, lycopene cyclase; crtX, β -carotene hydroxylase; crtZ, zeaxanthin glycosylase

Example 9 Construction of cDNA library of Phaffia rhodozyma

a) Isolation of total RNA from Phaffia rhodozyma

All solutions were made in DEPC-treated distilled water and all equipments were soaked overnight in 0.1% DEPC and then autoclaved.

A 300 ml Erlemeyer containing 60 ml YePD culture medium was inoculated with *Phaffia rhodozyma* strain CBS6938/1043-3 from a preculture to a final OD₆₀₀ of 0.1. This culture was incubated at 21 °C (300 rpm) until the OD₆₀₀ had reached 3-4.

The cells were harvest by centrifugation (4 °C, 8000 rpm, 5 min) and were resuspended in 12 ml of ice-cold extraction-buffer (0.1 M Tris-HCl, pH 7.5; 0.1 M LiCl; 0.1 mM EDTA). After centrifugation cells were resuspended in 2 ml of ice-cold extraction-buffer, 4 g of glassbeads (0.25 mm) and 2 ml phenol were added.

The mixture was vortexed 5 times at maximum speed for 30 s with 30 s cooling incubation intervals on ice.

The cell/glassbeads/phenol mixture was centrifuged (5 min, 15.300 rpm, 4 °C) and the aqueous phase (sup 1) was transferred to a fresh tube and was kept on ice.

The phenolic phase was retracted by adding an additional volume of 1 ml extraction buffer and 2 ml phenol.

After centrifugation (5 min, 15.300 rpm, 4 °C), the aquaous phase was transferred to sup 1 and extracted with an equal volume phenol:chloroform.

After centrifugation (5 min, 15.300 rpm, 4 °C), the aquaous phase was transferred to a fresh tube and 0.1 volume of 3 M NaAc; pH5.5 and 2.5 volumes of EtOH was added to precipitate RNA (incubation overnight -20 °C).

The precipitate was collected by centrifugation (10 min, 15.300 rpm, 4 °C) and drained off excess liquid and the RNA pellet was washed with 70 % icecold EtOH.

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After removing excess liquid the RNA was resuspended in 200 - 800 µl DEPC-treated water. RNA was stored at -70 °C. A 60 ml culture yielded 400 - 1500 µg total RNA. The integrity of total RNA was checked by formaldehyde RNA gel electrophoresis.

b) Selection of poly(A)* RNA

Isolation of poly(A)* from total RNA was carried out essential as described by Sambrook et al., 1989 (Molecular cloning, a laboratory manual, second edition) using the following solutions.

All solutions were prepared in DEPC-treated water and autoclaved.

10 RNA denaturation buffer:

1 M NaCl; 18% (v/v) DMSO.

Column-loading buffer (HEND): 10 mM Hepes, pH 7.6; 1 mM EDTA; 0.5 M Na Cl; 9% (v/v) DMSO.

Elution buffer (HE):

10 mM Hepes, pH 7.6; 1 mM EDTA.

Oligo(dT)-cellulose Type 7 was supplied by Pharmacia Biotech. O.1 g (dry weight) of oligo(dT)-cellulose was add to 1 ml HEND and the suspension was gently shaked for 1 h at 4 °C. Total RNA (1.5 mg dissolved in 500 µl) and 1 ml 1 M NaCl; 18% (v/v) DMSO was heated to 65 °C for 5 min. Then 600 µl NaCl/DMSO was added to the RNA, mixed and placed on ice for 5 min. The poly(A)* isolation was carried out be two cycles of purification. The final yield was about 45 µg poly(A)* RNA.

c) cDNA synthesis

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cDNAs were synthesized from 7.5 µg poly(A)*-RNAs using the cDNA Synthesis Kit (#200401; Strategene^R). Synthesis was carried out according to the instruction manual with some minor modification.

SuperScript[™] II RNase H⁻ Reverse Transcriptase (Gibco BRL) was used in the first strand reaction instead of MMLV-RT.

The following reagents were add in a microcentrifuge:

3 µl of poly(A)' RNAs

2 µl of linker-primer

23.5 µl DMQ

Incubate 10 min 70 °C, spin quickly in microcentrifuge and add,

10 µl of 5 x First Strand Buffer (provided by Gibco BRL)

5 μl of 0.1 M DTT (provided by Gibco BRL)

3 µl of first strand methyl nucleotide mixture

1 μl of RNase Block Ribonuclease Inhibitor (40 U/μl)

Annealling of template and primers by incubation the mixture at 25 °C for 10 min followed by 2 min at 42 °C and finally add;

2.5 ul SuperScript™ Il RNase H Reverse Transcriptase

First-strand reaction was carried out at 42 °C for 1 h.

Size fractionation was carried out using Geneclean^R II kit (supplied BIO 101, Inc.). The volume of the cDNA mixture obtained after Xhol digestion was brought up by adding DMQ to a final volume of 200 µl. Three volumes of NaI was added and the microcentrifuge tube was placed on ice for 5 min. The pellet of glassmilk was washed three times using 500 µl New Wash. Finally the cDNA was eluted in 20 µl DMQ.

The yield of cDNA was about 1 µg using these conditions.

d) cDNA cloning

cDNA library was constructed in the Uni-ZAPTM XR vector using 100 ng cDNAs. Ligation was performed two times overnight incubation at 12 °C. The cDNA library was packaged using the Packagene^R lambda DNA packaging system (Promega) according to the instruction manual. The calculated titer of the cDNA library was 3.5 10° pfu.

15 e) Mass excission

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Mass excision was carried out described in the protocol using derivatives of *E. coli* XL-Blue-MRF' as acceptor strain (see Table 5). Dilution of cell mixtures were plated onto 145 mm LB agar plates containing ampicillin, chloramphenicol and IPTG, yielding 250 - 7000 colonies on each plate. The plates were incubated overnight at 37 °C and further incubated one or two more days at room temperature.

Example 10

Cloning of the geranylgeranyl pyrophosphate synthase gene (crtE) of Phaffia rhodozyma

a) <u>lsolation of cDNA clone</u>

The entire library was excised into a farnesylpyrophosphate/ isopentenyl pyrophosphate accumulating cells of *E.coli* XL-Blue-MRF, which carries the plasmid pACCAR25ΔcrtE (further indicated as XL-Blue-MRF'[pACCAR25ΔcrtE]). The screening for the *crtE* gene was based on the color of the transformants. Introduction of the *crtB* gene in a genetic background of XL-Blue-MRF'[pACCAR25ΔcrtE] would result in a restoration of the complete route for the biosynthesis of zeaxanthin-diglucoside, which could be monitored by the presence of a yellow/orange pigmented colony. About 8.000 colonies were spread on LB agar plates containing appropriate antibiotics and IPTG. One colonie was found to have changed to a yellow/orange color.

b) Characterization of complementing cDNA clone

These colonies were streaked on LB-ampicillin agar plates. Plasmid DNA was isolated from this yellow colonies and found to include a 1.85 kb fragment (Fig 2A). The resulting plasmid, designated pPRcrtE,

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was used for retransformation experiments (Table 6). Only the transformation of XL-Blue-MRF'[pACCAR25\(\trice{L}\)crtE] with pPRcrtE resulted in a white to yellow color change in phenotype. To test whether the color change was due to complemention and not caused by cDNA alone pPRcrtE was transformed into XL-Blue-MRF'. Selection of transformants on LB-ampicillin agar plate containing IPTG did not result in color changes of the colonies (Table 6). Therefore we tentatively concluded, that we have cloned a cDNA of *P. rhodozyma* encoding GPPP synthase which is involved in the conversion of IPP and FPP to GGPP.

Table 6: Color phenotype of carotenoid producing E. coli strains transformed with pPRcrtE.

	pUC19 (control)	pPRcrtE
XL-Blue-MRF' (Ap. IPTG)	white	white
XL-Blue-MRF' [pACCAR25ΔcrtE] (Ap. Cm. IPTG)	white	yellow/orange
XL-Blue-MRF' [pACCAR25ΔcrtB] (Ap, Cm, IPTG)	white	white

Transformation: 10 ng of each plasmid was mixed to CaCl₂ competent E. coli cells. Transforment cells were selected by plating 1/10 and 1/100 volume of the DNA/cell mixture on LB agar-medium containing the appropriate antibiotics (in brackets).

c) Sequence analysis of cDNA fragment

Plasmid pPRcrtE was used to determine the nucleotide sequence of the 1.85 kb cDNA.

The sequence comprised 1830 nucleotides and a 31 bp poly(A) tail. An open reading frame (ORF) of 375 amino acids was predicted. The nucleotide sequence and deduced amino acid sequence are shown as SEQIDNO: NO 14 and 15, respectively. A search in SWISS-PROT protein sequence data bases using the Blitz amino acid sequence alignment program indicated amino acid homology (52 % in 132 aa overlap; *Neurospora crassa*) especially to the conserved domain I in geranylgeranyl-PPi synthase enzymes of different organisms (Botella et al., Eur. J. Biochem. (1995) 233; 238-248).

Example 11

Cloning of the phytoene synthase gene (crtB) of Phaffia rhodozyma

a) Isolation of cDNA clone

The entire library was excised into a geranylgeranylpyrophosphate accumulating cells of *E.coli* XL-Blue-MRF', which carries the plasmid pACCAR25ΔcrtB (further indicated as XL-Blue-MRF'[pACCAR25ΔcrtB]). The screening for the *crtB* gene was based on the color of the transformants.

Introduction of the *crtB* gene in a genetic background of XL-Blue-MRF'[pACCAR25\(\triangle\) crtB] would result in a restoration of the complete route for the biosynthesis of zeaxanthin-diglucoside, which could be monitored by the presence of a yellow/orange pigmented colony.

About 25.000 colonies were incubated on LB agar plates containing appropriate antibiotics and IPTG.

Three colonies were found to have changed to a yellow/orange color.

b) Characterization of complementing cDNA clone

These colonies were streaked on LB-ampicillin agar plates. Plasmid DNA. designated pPRcrtB1 to 3, was isolated from these yellow colonies and found to include a 2.5 kb fragment (Fig 2B). One of the resulting plasmids, pPRcrtB1 was used for retransformation experiments (Table 7). Only the transformation of XL-Blue-MRF'[pACCAR25\Delta crtB] with pPRcrtB resulted in a white to yellow color change in phenotype. Therefore we tentative conclude that we have cloned a cDNA of *P. rhodozyma* encoding phytoene synthase which is involved in the conversion of 2 GGPP molecules via prephytoene pyrophosphate into phytoene.

Table 7: Color phenotype of carotenoid producing E. coli strains transformed with pPRcrtB.

	pUC19 (control)	pPRcrtB
XL-Blue-MRF' (Ap, IPTG)	white	white
XL-Blue-MRF' [pACCAR25\(\Delta\text{pr}\) (Ap, Cm, IPTG)	white	yellow/orange
XL-Blue-MRF' [pACCAR25ΔcπΕ (Ap, Cm, IPTG)	white	white

Legend: see Table 6.

c) Sequence analysis of cDNA fragment.

Plasmid pPRcrtB2, which contains the longest cDNA insert, was used to determine the nucleotide sequence of the 2.5 kb cDNA. The sequence comprised 2483 nucleotides and a 20 bp poly(A) tail. An open reading frame (ORF) of 684 amino acids was predicted. The nucleotide sequence and deduced amino acid sequence are shown in SEQIDNOs: 12 and 13, respectively. A search in SWISS-PROT protein sequence data bases using the Blitz amino acid sequence alignment program Data indicated some amino acid homology (26 % identity in 441 aa overlap of crtB gene of Neurospora crassa) with crtB genes of other organisms.

Example 12

Cloning of the phytoene desaturase gene (crt1) of Phaffia rhodozyma

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a) Isolation of cDNA clone

The entire library was excised into a phytoene accumulating cells of *E.coli* XL-Blue-MRF', which carries the plasmid pACCAR25\(\Delta\crit{Ctt}\) (further indicated as XL-Blue-MRF'[pACCAR25\(\Delta\crit{Ctt}\)]). The screening for the *crtl* gene was based on the color of the transformants. Introduction of the *crtl* gene in a genetic background of XL-Blue-MRF'[pACCAR25\(\Delta\crit{Ctt}\)] would result in a restoration of the complete route for the biosynthesis of zeaxanthin, which could be monitored by the presence of a yellow/orange pigmented colony.

About 14.000 colonies were incubated on LB agar plates containing appropriate antibiotics and IPTG. Two colonies were found to have changed to a yellow/orange color.

b) Characterization of complementing cDNA clones

These colonies were streaked on LB-ampicillin agar plates. Plasmid DNA, designated pPRcrtl.1 and pPRcrtl.2, was isolated from these yellow colonies and found to include a 2.0 kb fragment (Fig 2C). One of the resulting plasmids, pPRcrtl.1 was used for retransformation experiments (Table 8). Only the transformation of XL-Blue-MRF'[pACCAR25 Δ crtX Δ crtl] with pPRcrtl resulted in a white to yellow color change in phenotype. Therefore we tentative conclude that we have cloned a cDNA of P. rhodozyma encoding phytoene desaturase which is involved in the conversion of phytoene to lycopene.

Table 8: Color phenotype of carotenoid producing E. coli strains transformed with pPRcrt1.

	pUC19	pPRcni
XL-Blue-MRF' (Ap, IPTG)	white	white
XL-Blue-MRF' [pACCAR25ΔcrtX Δcrtl (Ap, Cm, IPTG)	white	yellow/orange
XL-Blue-MRF' [pACCAR25ΔcrtB (Ap, Cm, IPTG)	white	white

Legend: see Table 6.

c) Sequence analysis of cDNA fragment

One of the plasmid pPRcrtI was used to determine the nucleotide sequence of the 2.0 kb cDNA. The sequence comprised 2038 nucleotides and a 20 bp poly(A) tail. An open reading frame (ORF) of 582 amino acids was predicted. The nucleotide sequence and deduced amino acid sequence are shown in SEQIDNOs: 16 and 17, respectively. A search in SWISS-PROT protein sequence data bases using the Blitz amino acid sequence alignment program Data indicated amino acid homology to phytoene desaturase gene of *N. crassa* (53% identity in 529 aa overlap).

Example 13 Cloning of the lycopene cyclase gene (crtY) of Phaffia rhodozyma

a) Isolation of cDNA clone

The entire library was excised into a lycopene accumulating cells of *E.coli* XL-Blue-MRF', which carries the plasmid pACCRT-EIB (further indicated as XL-Blue-MRF'[pACCRT-EIB]). The screening for the *crtY* gene was based on the color of the transformants. Introduction of the *crtY* gene in a genetic background of XL-Blue-MRF'[pACCRT-EIB] would result in a restoration of the complete route for the biosynthesis of β-carotene, which could be monitored by the presence of a yellow pigmented colony.

About 8.000 colonies were incubated on LB agar plates containing appropriate antibiotics and IPTG.

One colony was found to have changed to a yellow color.

b) Characterization of complementing cDNA clone

This colony was streaked on LB-ampicillin agar plates. Plasmid DNA was isolated from this yellow colony and found to include a 2.5 kb fragment (Fig 2B). The resulting plasmid, designated pPRcrtY, was used for retransformation experiments (Table 9. Surprisingly, not only transformation of XL-Blue-MRF'[pACCRT-EIB] but also transformation of XL-Blue-MRF'[pACCAR25ΔcrtB] with pPRcrtY resulted in a red to yellow color change in phenotype.

Table 9: Color phenotype of carotenoid producing E. coli strains transformed with pPRcrtY.

	pUC19	pPRcrtB	
XL-Blue-MRF' (Ap, IPTG)	white	white	
XL-Blue-MRF' [pACCRT-EIB (Ap, Cm, IPTG)	red	yellow	
XL-Blue-MRF' [pACCAR25∆crtB (Ap, Cm, IPTG)	red	yellow	

Legend: see Table 6.

A second transformation experiment was carried out including the previously cloned cDNA of pPRcrtB. As shown in table 6 the cDNA previously (example 3) isolated as encoding phytoene synthase was able to complement the crtY deletion resulting in the biosynthesis of β-carotene in XL-Blue-MRF'[pACCRT-EIB].

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Sequence analysis of the cDNA insert of pPRcrtY (SEQIDNOs: 18 and 19) showed that it was similar to the sequence of cDNA fragment of pPRcrtB.

From these data we tentative conclude that we have cloned a cDNA of P. rhodozyma encoding phytoene synthase and lycopene cyclase which is involved in the conversion of 2 GGPP molecules via prephytoene pyrophosphate into phytoene and lycopene to β -carotene, respectively. This is the first gene in a biosynthetic pathway of carotenoids synthesis that encodes two enzymatic activities.

Table 10: Color phenotype of carotenoid producing *E. coli* strains transformed with different cDNAs of *Phaffia rhodozyma* (Ap, Cm, IPTG).

	pUC19	pPRcrtE	pPRcrtB	pPRcrtY
XL-Blue-MRF' [pACCAR25ΔcπE]	white	yellow/ orange	white	white
XL-Blue-MRF' [pACCAR25ΔcπB]	white	white	yellow/ orange	yellow/ orange
XL-Blue-MRF' [pACCRT-EIB]	red	red	yellow	yellow

Legend: see Table 6

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Example 14

Cloning of the isopentenyl diphosphate (IPP) isomerase gene (idi) of Phaffia rhodozyma

a) Isolation of cDNA clone

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The entire *Phaffia* cDNA library was excised into lycopene accumulating cells of *E.coli* XL-Blue-MRF', each carrying the plasmid pACCRT-EIB (further indicated as XL-Blue-MRF'[pACCRT-EIB]).

About 15.000 colonies were incubated on LB agar plates containing appropriate antibiotics and IPTG. One colony was found to have a dark red colour phenotype.

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b) Characterization of complementing cDNA clone

This colony was streaked on LB-ampicillin agar plates. Plasmid DNA was isolated from this yellow colony and found to include a 1.1 kb fragment. The resulting plasmid, designated pPRcrtX, was used for retransformation experiments (Table 11).

All colonies of XL-Blue-MRF'[pACCAR-EIB] transformed with pPRcrtX had a dark red phenotype. From these data we tentatively concluded, that we have cloned a cDNA of *P. rhodozyma* expression of which results in an increased lycopene production in a genetically engineered *E. coli* strain.

Table 11: Color phenotype of carotenoid producing E. coli strains transformed with pPRcrtX.

	pUC19	pPRcrtX	
XL-Blue-MRF' (Ap, IPTG)	white	white	
XL-Blue-MRF' [pACCRT-EIB (Ap, Cm, IPTG)	red	dark red	

Legend: see Table 6.

c) Sequence analysis of cDNA fragment

In order to resolve the nature of this gene the complete nucleotide sequence of the cDNA insert in pPRcrtX was determined. The nucleotide sequence consist of the 1144 bp. The sequence comprised 1126 nucleotides and a poly(A) tail of 18 nucleotides. An open reading frame (ORF) of 251 aminoacids with a molecular mass of 28.7 kDa was predicted. The nucleotide sequence and deduced amino acid sequence are shown in SEQIDNOs: 20 and 21, respectively.

A search in SWISS-PROT protein sequence data bases using the Blitz amino acid sequence alignment program Data indicated aminoacid homology to isopentenyldiphosphate (IPP) isomerase (idi) of S. cerevisiae (42.2 % identity in 200 aminoacid overlap). IPP isomerase catalyzes an essential activation step in the isoprene biosynthetic pathway which synthesis the 5-carbon building block of carotenoids. In analogy to yeast the gene of *Phaffia* was called idi1. The cDNA clone carrying the genes was then called pPRidi.

Example 15

Overexpression of the idi gene of P. rhodozyma in a carotenogenic E. coli

Lycopene accumulating cells of *E.coli* XL-Blue-MRF', which carry the plasmid pACCRT-EIB (further indicated as XL-Blue-MRF'[pACCRT-EIB]) were transformed with pUC19 and pPRidi and transformants were selected on solified LB-medium containing Amp and Cm. The transformants, called XL-Blue-MRF'[pACCRT-EIB/pUC19 and [pACCRT-EIB/pPRidi], were cultivated in 30 ml LB-medium containing Amp, Cm and IPTG at 37 °C at 250 rpm for 16 h. From these cultures 1 ml was used for carotenoid extraction and analysis. After centrifugation the cell pellet was dissolved in 200 μ l aceton and incubated at 65 °C for 30 minutes. Fifty μ l of the cell-free aceton fraction was then used for high-performance liquid chromatography (HPLC) analysis. The column (chrompack cat. 28265; packing nucleosil 100C18) was developed with water-acetonitrile-2-propanol (from 0 to 45 minutes 9:10:8) and after 45 minutes 2:18:80) at a flow rate of 0.4 ml per minute and recorded with a photodiode array detector at 470 +/- 20 nm. Lycopene was shown to have a retention time of about 23 minutes under these conditions. The peak area was used as the relative lycopene production (mAu*s). The relative

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lycopene production was 395 and 1165 for XL-Blue-MRF'[pACCRT-EIB/pUC19] and [pACCRT-EIB/pPRidi], respectively.

These data show the potentials of metabolic pathway engineering in *Phaffia*, as increased expression of the *idi* of *Phaffia rhodozyma* causes a 3-fold increase in carotenoid biosynthesis in *E. coli*.

This cDNA may be over-expressed in a transformed *Phaffia* cell with a view to enhance carotenoid and/or xanthophyll levels. The cDNA is suitably cloned under the control of a promoter active in *Phaffia*, such as a strong promoter according to his invention, for example a *Phaffia* glykolytic pathway promoter, such as the GAPDH-gene promoter disclosed herein, or a *Phaffia* ribosomal protein gene promoter according to the invention (*vide sub*). Optionally, the cDNA is cloned in front of a transcriptional terminator and/or polyadenylation site according to the invention, such as the GAPDH-gene terminator/polyadenylation site. The feasibility of this approach is illustrated in the next example, where the *crt*B gene from *Erwinia uredovora* is over-expressed in *Phaffia rhodozyma* by way of illustration.

Example 16

Heterologous expression of carotenogenic gene from Erwinia uredovora in Phaffia rhodozyma.

The coding sequence encoding phytoene synthase (crtB) of Erwinia uredovora (Misawa et al., 1990) was cloned between the promoter and terminator sequences of the gpd (GAPDH-gene) of Phaffia by fusion PCR. In two separate PCR reactions the promoter sequence of gpd and the coding sequence of crtB were amplified. The former sequence was amplified using the primers 5177 and 5128 and pPR8 as template. This latter vector is a derivative of the Phaffia transformation vector pPR2 in which the promoter sequence has been enlarged and the Bg/II restriction site has been removed. The promoter sequence of gpd was amplified by PCR using the primers 5226 and 5307 and plasmid pPRgpd6 as template. The amplified promoter fragment was isolated, digested with KpnI and BamHI and cloned in the Kpnl-BglII fragment of vector pPR2, yielding pPR8. The coding sequence of crtB was amplified using the primers 5131 and 5134 and pACCRT-EIB as template. In a second fusion PCR reaction, using the primers 5177 and 5134, 1 μ g of the amplified promoter and crtB coding region fragment used as template yielding the fusion product Pgpd-crtB. The terminator sequence was amplified under standard PCR conditions using the primers 5137 and 5138 and the plasmid pPRgdh6 as template. Primer 5137 contains at the 5' end the last 11 nucleotides of the coding region of the crtB gene of E. uredovora and the first 16 nucleotides of the terminator sequence of gpd gene of P. rhodozyma. By a two basepair substitution a BamH1 restriction site was introduced. The amplified fusion product (Pgpd-crtB) and the amplified terminator fragments were purified and digested with HindIII and BamHI and cloned in the dephosphorylated HindIII site of the cloning vector pMTL25. The vector with the construct Pgpd-crtB-Tgpd was named pPREX1.1.

The *Hind*III fragment containing the expression cassette Pgpd-crtB-Tgpd was isolated from pPREX1.1 and ligated in the dephosphorylated *Hind*III site of the *Phaffia* transformation vector pPR8. After transformation of the ligation mixture into *E. coli* a vector (pPR8crtB6.1) with the correct insert was chosen for *Phaffia* transformation experiments.

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Phaffia strain CBS6938 was transformed with pPR8crtB6.1, carrying the expression cassette Pgpd-crtB-Tgpd, and transformants were selected on plates containing G418. The relative amount of astaxanthin per OD₆₆₀ in three G418-resistant transformants and the wild-type Phaffia strains was determined by HPLC analysis (Table 12). For carotenoid isolation from Phaffia the method of DMSO/hexane extraction described by Sedmak et al., (1990; Biotechn. Techniq. 4, 107-112) was used.

Table 12. The relative astaxanthin production in a *Phaffia* transformant carrying the crtB gene of E. uredovora.

	of astaxanthin	
Strain:	(mAU*s/OD ₆₆₀)	
P. rhodozyma CBS6938	448	
P. rhodozyma CBS6938		
[pPR8cr1B6.1]#1	626	
[pPR8crtB6.1]#2	716	
[pPR8crtB6.1]#4	726	

5128: 5' caactgccatgatggtaagagtgttagag 3'

5177: 5' cccaagetttctcgaggtacctggtgggtgcatgtatgtac3'

5131: 5' taccatcatggcagttggctcgaaaag 3'

5134: 5' cccaagcttggatccgtctagagcggggggcgctgcc3'

5137: 5' ccaaggeetaaaeggateeeteeaaace 3'

5138: 5' gccaagettctcgagettgatcagataaagatagagat3'

5307: 5' gttgaagaagggatccttgtggatga 3'

The gpd sequences are indicated in bold, the crtB sequences in italic, additional restriction sites for cloning are underlined and base substitution are indicated by double underlining.

Example 17

Isolation and characterization of the crtB gene of Phaffia

It will also be possible to express the *Phaffia rhodozyma* gene corresponding to *crt*B and express it under the control of its own regulatory regions, or under the control of a promoter of a highly expressed gene according of the invention. The *Phaffia* transformation procedure disclosed herein, invariably leads to stably integrated high copy numbers of the introduced DNA, and it is expected, that expression of the gene under the control of its own promoter will also lead to enhanced production of

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carotenoids, including astaxanthin. To illustrate the principle, a protocol is given for the cloning of the crtB genomic sequence, below.

To obtain the genomic criB-gene including expression signals the 2.5 kb BamHI-XhoI fragment was isolated from the vector pPRcrtB and used as probe to screen a cosmid library of Phaffia.

The construction and screening of the library was carried out as described in Example 3 using the *crtB* gene as probe instead of the *gapdh*-gene.

After the rounds of hybridization, 2 colonies were identified giving a strong hybridization signal on the autoradiogram after exposure. Cosmid DNA isolated from these colonies was called pPRgcrtB#1.1 and pPRgcrtB#7, respectively.

Chromosomal DNA isolated from Phaffia rhodozyma strain CBS 6938 and cosmid pPRgcrtB#7 was digested with several restriction enzymes. The DNA fragments were separated, blotted and hybridized with a amino-terminal specific probe (0.45 kb Xbal fragment) of crtB under conditions as described before. After exposure, the autoradiogram showed DNA fragments of different length digested by different restriction enzymes which hybridized with the crtB probe. On the basis that no EcoRI site is present in the cDNA clone a EcoRI fragment of about 4.5 kb was chosen for subcloning experiments in order to determine the sequence in the promoter region and to establish the presence of intron sequences in the crtB gene. A similar sized hybridizing fragment was also found in the chromosomal DNA digested with EcoRI. The fragment was isolated from an agarose gel and ligated into the corresponding site of pUC19. The ligation mixture was transformed to competent E. coli cells. Plasmids with the correct insert in both orientations, named pPR10.1 and pPR10.2, were isolated from the transformants. Comparison of the restriction patterns of pPR10.1/pPR10.2 and pPRcrtB digested with Xbal gave an indication for the presence of one or more introns as the internal 2.0 kb Xbal fragment in the cDNA clone was found to be larger in the former vectors. The subclone pPR10.1 was used for sequence analysis of the promoter region and the structural gene by the so-called primer walking approach. The partial sequence of the insert in show in SEQIDNO: 22. Comparison of the cDNA and the genomic sequence revealed the presence of 4 introns.

Example 18

Isolation of promoter sequences with high expression levels

This example illustrates the the feasibility of the "cDNA sequencing method" referred to in the detailed description, in order to obtain transcription promoters from highly expressed genes.

For the isolation and identification of transcription promoter sequences from *Phaffia rhodozyma* genes exhibiting high expression levels, the cDNA library of *Phaffia rhodozyma* was analyzed by the following procedure.

The cDNA library was plated on solified LB-medium containing Amp and 96 colonies were randomly picked for plasmid isolation. The purified plasmid was digested with Xhol and Xbal and loaded on a agarose gel. The size of the cDNA inserts varied from 0.5 to 3.0 kb. Subsequently, these plasmids were used as template for a single sequence reaction using the T3 primer. For 17 cDNA clones no sequence data were obtained. The sequences obtained were translated in all three reading frames. For

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each cDNA sequence the longest deduced amino acid sequences were compared with the SwissProt protein database at EBI using the Blitz program. For 18 deduced amino acid sequences no homology to known proteins was found whereas six amino acid sequences showed significant homology to hypothetical proteins. Fifty-five amino acid sequences were found to have significant homology to proteins for which the function is known. About 50 % (38/79) were found to encode ribosomal proteins of which 12 full-length sequences were obtained.

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Table 13. Overview of expressed cDNAs, encoded proteins and reference to the Sequence Listing

cDNA	coding for	SEQIDNO
10	ubiquitin-40S	24
11	Glu-repr.gene	26
18	. 40S rib.prot S27	28
35	60S rib.prot PIα	30
38	60S rib.prot L37e	32
46	60S rib.prot L27a	34
64	60S rib.prot L25	36
68	60S rib.prot P2	38
73	40S rib.prot S17A/B	40
76	40S rib.prot S31	42
78	40s rib.prot S10	44
85	60S rib.prot L37A	46
87	60S rib.prot L34	48
95	60S rib.prot S16	50

By sequence homology it was concluded that in *Phaffia* the 40S ribisomal protein S37 is fused to ubiquitin as is found in other organisms as well. The nucleotide sequences and deduced amino acid sequences of the full length cDNA clones are listed in the sequence listing. Six ribosomal proteins were represented in the random pool by more than one individual cDNA clone. The 40S ribosomal proteins S10 (SEQIDNO:44), S37 (+ ubiquitin) (SEQIDNO:24) and S27 (SEQIDNO:28) were represented twice and 60S (acidic) ribosomal proteins P2 (SEQIDNO:38), L37 (SEQIDNO:46) and L25 (SEQIDNO:36) found three times. From these results we conclude, that these proteins are encoded by multiple genes or that these genes are highly expressed. Therefore isolation of these promoter sequences are new and promissing target sequences to isolate high level expression signals from *Phaffia rhodozyma*. Furthermore, a cDNA clone was isolated which showed 50 % homology to an abundant glucose-repressible gene from *Neurospora crassa* (Curr. genet. 14: 545-551 (1988)). The nucleotide sequence and the deduced amino acid sequence is shown in SEQIDNO:26. One of the advantages of such a promoter sequence is that it can be used to separated growth (biomass accumulation) and gene expression (product accumulation) in large scale *Phaffia* fermentation.

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For the isolation of the promoter sequences of interest (as outlined above) a fragment from the corresponding cDNA clone can be used as probe to screen the genomic library of *Phaffia rhodozyma* following the approach as described for the GAPDH-gene promoter (Example 3, *supra*). Based on the determined nucleotide sequence of the promoter, specific oligonucleotides can be designed to construct a transcription fusion between the promoter and any gene of interest by the fusion PCR technique, following the procedure as outlined in Example 5 (*supra*).

Example 19

Isolation of carotenogenic genes by heterologous hybridization

For the identification and isolation of corresponding carotenoid biosynthetic pathway genes from organisms related to *Phaffia rhodozyma* heterologous hybridization experiments were carried out under conditions of moderate stringency. In these experiments chromosomal DNA from two carotenogenic fungi (*Neurospora crassa* and *Blakeslea trispora*) and the yeasts *S. cerevisiae* and three yeast and fungal species from the genus *Cystofylobasidium* was used. These three carotenogenic yeasts are, based on phylogenetic studies, the ones most related to *P. rhodozyma*.

Chromosomal DNA from the yeast species Cystofylobasidium infirmo-miniatum (CBS 323), C. bisporidii (CBS 6346) and C. capitatum (CBS 6358) was isolated according the method as developed for Phaffia rhodozyma, described in example 3 of European patent application 0 590 707 A1; the relevant portions of which herein incorporated by reference. Isolation of chromosomal DNA from the fungi Neurospora crassa and Blakeslea trispora was essentially carried as described by Kolar et al. (Gene, 62: 127-134), the relevant parts of which are herein incorporated by reference.

Chromosomal DNA (5 µg) of C. infirmo-miniatum, C. bisporidii, C. capitatum, S. cerevisiae, P. rhodozyma, N. crassa and B. trispora was digested using EcoRI. The DNA fragments were separated on a 0.8% agarose gel, blotted and hybridized using the following conditions.

Hybridization was carried out at two temperatures (50 °C and 55 °C) using four different ³²P labelled *Phaffia* probes. The probes were made using random primed hexanucleotide labellings reactions using the *Xhol-Xbal* fragment(s) from the cDNA clones pPRcrtE, pPRcrtB, pPRcrtI and pPRidi as template. Hybridization was carried out o/n (16 h) at the indicated temperatures. After hybridization the filters were washed 2 times for 30 min. at the hybridization temperatures using a solution of 3*SSC; 0.1 % SDS; 0.05% sodiumpyrophosphate. Films were developed after exposure of the filters to X-ray films in a cassette at -80 °C for 20 h.

Using the cDNA clone of crtE of P. rhodozyma faint signals were obtained for C. infirmominiatum, C. capitatum. Using the cDNA clone of crtB of P. rhodozyma strong signals were obtained to the high molecular weight portion of DNA from C. infirmo-miniatum and C. capitatum. Furthermore a strong signal was obtained in the lane loaded with digested chromosomal DNA from B. trispora. Only a faint signal was obtained for C. capitatum at 50 °C using the cDNA clone of crtl of P. rhodozyma. Using the cDNA clone of idi of P. rhodozyma faint signals were obtained with chromosomal DNA from C. infirmo-miniatum, C. bisporidii and C. capitatum at both temperatures. A strong signal was obtained in the lane loaded with digested chromosomal DNA from B. trispora.

We conclude, that carotenoid biosynthesis cDNAs or genes, or *idi* cDNAs or genes, can be isolated from other organisms, in particular from other yeast species by cross-hybridisation with the cDNA fragments coding for *P. rhodozyma* carotenoid biosynthesis enzymes, or isopentenyl pyrophosphate isomerase coding sequences respectively, using moderately stringent hybridisation and washing conditions (50 °C to 55 °C, 3xSSC).

Deposited microorganisms

E. coli containing pGB-Ph9 has been deposited at the Centraal Bureau voor Schimmelcultures, Oosterstraat 1, Baarn, The Netherlands, on June 23, 1993, under accession number CBS 359.3.

The following strains have been deposited under the Budapest Treaty at the Centraal Bureau voor Schimmelcultures, Oosterstraat I, Baarn, The Netherlands, on February 26, 1996:

ID nr.	Organism	relevant feature	Deposit number
DS31855	E. coli	crtY of P. rhodozyma	CBS 232.96
DS31856	E. coli	crtl of P. rhodozyma	CBS 233.96
DS31857	E. coli	crtE of P. rhodozyma	CBS 234.96
DS31858	E. coli	crtB of P. rhodozyma	CBS 235.96

SEQUENCE LISTING

	(1) GENERAL INFORMATION:	
5	(i) APPLICANT: (A) NAME: Gist-brocades B.V.	
10	(B) STREET: Wateringseweg l (C) CITY: Delft (E) COLIMIRY: The Netherlands (F) POSTAL CODE (ZIP): 2611 XT	
	(ii) TITLE OF INVENTION: Improved methods for transforming Phaffia a recombinant INA for use therein	nd
15	(iii) NUMBER OF SEQUENCES: 51	
20	(iv) COMPUTER READABLE FORM: (A) MEDIUM TYPE: Floppy disk (B) COMPUTER: IBM PC compatible (C) OPERATING SYSTEM: PC-DOS/MS-DOS (D) SOFTWARE: Patentin Release #1.0, Version #1.25 (EPO)	
ಚ	(v) CURRENT APPLICATION DATA: APPLICATION NUMBER:	
	(2) INFORMATION FOR SEQ ID NO:1:	
30	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 25 base pairs (B) TYPE: nucleic acid (C) STRANDEINESS: single (D) TOPOLOGY: linear	
35	(ii) MOLECULE TYPE: DNA (genomic)	
	(iii) HYPOTHETICAL: NO	
40	(vi) ORIGINAL SOURCE: (C) INDIVIDUAL ISOLATE: AB3005	
	(xa) SEQUENCE DESCRIPTION: SEQ ID NO:1:	
45	COOGATCCAA RCINACNOCN AIGGC	25
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55	(ii) MOLECULE TYPE: DNA (genomic)	
	(iii) HYPOTHETICAL: NO	
60	(vi) ORIGINAL SOURCE: (C) INDIVIDUAL ISOLATE: AB3006	
65	<pre>(ix) FEATURE: (A) NAME/KEY: misc_feature (B) LOCATION: one-of(12) (D) OTHER INFORMATION: /note= "N at position 12 is inosine"</pre>	
70	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:	

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	(ii) MOLECULE TYPE: DNA (genomic)	
	(iii) HYPOTHETICAL: NO	
5	(vi) ORIGINAL SOURCE: (C) INDIVIDUAL ISOLATE: AB5177	
10	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:6:	
	CCCAAGCTIC TOGAGGIACC TOGIGOGIGC ATGIATGIAC	40
	(2) INFORMATION FOR SEQ ID NO:7:	
15	 (i) SEQUENCE CHARACTERISTICS: (A) LENGIH: 30 base pairs (B) TYPE: nucleic acid (C) STRANDEINESS: single (D) TOPOLOGY: linear 	
20	(ii) MOLECULE TYPE: DNA (genomic)	
-	(iii) HYPOTHETICAL: NO	
25	(vi) ORIGINAL SOURCE: (C) INDIVIDUAL ISOLATE: AB5137	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:7:	
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	(2) INFORMATION FOR SEQ ID NO:8:	
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40	(ii) MOLECULE TYPE: DNA (genomic)	
	(iii) HYPOTHETICAL: NO	•
45	(VI) ORIGINAL SOURCE: (C) INDIVIDUAL ISOLATE: AB5138	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:8:	
50	GCCAAGCTTC TOGAGCTTGA TCAGATAAAG ATAGAGAT	38
	(2) INFORMATION FOR SEQ ID NO:9:	
55	(i) SEQUENCE CHARACTERISTICS: (A) LENGIH: 2309 base pairs (B) TYPE: nucleic acid (C) STRANDEINESS: double (D) TOPOLOGY: linear	
64	(ii) MOLECULE TYPE: DNA (genomic)	
	(iii) HYPOTHETICAL: NO	
6	(iv) ANTI-SENSE: NO	
	(vi) ORIGINAL SOURCE:(A) ORGANISM: Phaffia rhodozyma(B) STRAIN: CBS 6938	
	70	

	(ix)	FEATURE: (A) NAME/KEY: excn (B) LOCATION: 300330	
5	(ix)	FEATURE: (A) NAME/KEY: intron (B) LOCATION: 331530	
10	(ix)	FEATURE: (A) NAME/KEY: excn (B) LOCATION: 531578	
15	(ix)	FEATURE: (A) NAME/KEY: intron (B) LOCATION: 579668	
20	(ix)	FEATURE: (A) NAME/KEY: exon (B) LOCATION: 669690	
20	(ix)	FEATURE: (A) NAME/KEY: intron (B) LOCATION: 691767	
25	(ix)	FEATURE: (A) NAME/KEY: excan (B) LOCATION: 768805	
30	(ix)	FEATURE: (A) NAME/KEY: intron (B) LOCATION: 806905	
35	(xi)	FEATURE: (A) NAME/KEY: excn (B) LOCATION: 906923	
	(ix)	FEATURE: (A) NAME/KEY: intron (B) LOCATION: 9241030	
40	(ix)	FEATURE: (A) NAME/KEY: exon (B) LOCATION: 10311378	
45	(ix)	FEATURE: (A) NAME/KEY: intron (B) LOCATION: 13791508	
50	(ix)	FEATURE: (A) NAME/KEY: exon (B) LOCATION: 15092020	
55	(ix)	FEATURE: (A) NAME/KEY: CDS (B) LOCATION: join(300330, 531578, 669690, 768805,923, 10311378, 15092020)	906
60		SEQUENCE DESCRIPTION: SEQ ID NO:9:	
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		SC GTCTCCCCCC GGCAACCCCC GGTGCCCCCCC TCCGCTTACG TCAGCCACCC	120
65		IC CATCICITIC TCICICCITC CAAAAGICHT TCAGHTHAA ACGGCCCCCA	180
		AG AGGGGACTIT TICTITICCIT CICCCCATCA TCCACAAAGA TCICTCITCI	240
70	1 CAACAAC	AA CIACIACIAC TACCACIACC ACCACIACIT CTCTAACACT CTTACCATC	299

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ATC CAC GAC AAC TAC ACC A GICAGICCTT INCITIIGGAC TIGICIGGCC

Ile His Asp Asn Tyr Thr 165

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	GAT GTC TCC GTC GTC GAT CTT GTC GTC CGA ATC GAG AAG GCC GCC TCT Asp Val Ser Val Val Asp Leu Val Val Arg Ile Glu Lys Gly Ala Ser 240 245 250	1762
30	TAC GAG GAG ATC AAG GAG ACC ATC AAG AAG GCC TCC CAG ACC CCT GAG Tyr Glu Glu Ile Lys Glu Thr Ile Lys Lys Ala Ser Gln Thr Pro Glu 255 260 265	1810
35	CTC AAG GGT ATC CTG AAC TAC ACC GAC GAC GAC GTC GTC TCC ACC GAT Leu Lys Gly Ile Leu Asn Tyr Thr Asp Asp Gln Val Val Ser Thr Asp 270 280 285	1858
40	TTC ACC GGT GAC TCT GCC TCC TCC ACC TTC GAC GCC CAG GGC GGT ATC Phe Thr Gly Asp Ser Ala Ser Ser Thr Phe Asp Ala Gln Gly Gly Ile 290 295 300	1906
45	TCC CTT AAC GGA AAC TTC GTC AAG CTT GTC TCC TGG TAC GAC AAC GAG Ser Leu Asn Gly Asn Phe Val Lys Leu Val Ser Trp Tyr Asp Asn Glu 305 310 315	1954
	TOG GCA TAC TOT GCC CGA GTC TGC GAC CIT GTT TCT TAC ATC GCC GCC Trp Gly Tyr Ser Ala Arg Val Cys Asp Leu Val Ser Tyr Ile Ala Ala 320 325 330	2002
50	CAG GAC GCC AAG GCC TAAAACGGFTC TCTCCAAACC CTCTCCCCTT TTGCCCTGCC Gln Asp Ala Lys Ala 335	2057
55	CATTGAATTG ATTCCCTAAA TAGAATATCC CACTTTCTTT TATGCTCTAC CTATGATCAG	2117
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	AAGGCAAGAA GTGAGCAAGA TATGAACAAG AACAACAAAG AAAAACAGAC AAAGAAAAAA	2237
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(2) INFORMATION FOR SEQ ID NO:10:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 338 amino acids
(B) TYPE: amino acid
(D) TOPOLOGY: linear

70

- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:10:
- Met Ala Val Lys Val Gly Ile Asn Gly Phe Gly Arg Ile Gly Arg Ile
 1
 5
 10
 15
 - Val Leu Arg Asn Ala Ile Ile His Gly Asp Ile Asp Val Val Ala Ile 20 25 30
- Asm Asp Pro Phe Ile Asp Leu Glu Tyr Met Val Tyr Met Phe Lys Tyr
- 15 Asp Ser Thr His Gly Val Phe Lys Gly Ser Val Glu Ile Lys Asp Gly 50 55 60
 - Lys Leu Val Ile Glu Gly Lys Pro Ile Val Val Tyr Gly Glu Arg Asp 65 70 75 80
- Pro Ala Asm Ile Gln Trp Gly Ala Ala Gly Ala Asp Tyr Val Val Glu 85 90 95
- Ser Thr Gly Val Phe Thr Thr Gln Glu Lys Ala Glu Leu His Leu Lys $_{\rm 25}$ $\,$ $\,$ 100 $\,$ $\,$ 105 $\,$ 110 $\,$
 - Gly Gly Ala Lys Lys Val Val Ile Ser Ala Pro Ser Ala Asp Ala Pro 115 120 125
- Met Phe Val Cys Gly Val Asn Leu Asp Lys Tyr Asp Pro Lys Tyr Thr 130 135 140
 - Val Val Ser Asn Ala Ser Cys Thr Thr Asn Cys Leu Ala Pro Leu Gly 145 150 155 160
- Lys Val Ile His Asp Asn Tyr Thr Ile Val Glu Gly Leu Met Thr Thr 165 170 175
- Val His Ala Thr Thr Ala Thr Gln Lys Thr Val Asp Gly Pro Ser Asn 40 180 185 190
 - Lys Asp Trp Arg Gly Gly Arg Gly Ala Gly Ala Asn Ile Ile Pro Ser 195 200 205
- 45 Ser Thr Gly Ala Ala Lys Ala Val Gly Lys Val Ile Pro Ser Leu Asn 210 215 220
 - Gly Lys Leu Thr Gly Met Ala Phe Arg Val Pro Thr Pro Asp Val Ser 225 230 235 240
 - Val Val Asp Leu Val Val Arg Ile Glu Lys Gly Ala Ser Tyr Glu Glu 245 250 255
- Ile Lys Glu Thr Ile Lys Lys Ala Ser Gln Thr Pro Glu Leu Lys Gly 260 265 270
 - Ile Leu Asn Tyr Thr Asp Asp Gln Val Val Ser Thr Asp Phe Thr Gly 275 280 285
- Asp Ser Ala Ser Ser Thr Phe Asp Ala Gln Gly Gly Ile Ser Leu Asn 290 295 300
- Gly Asn Phe Val Lys Leu Val Ser Trp Tyr Asp Asn Glu Trp Gly Tyr 305 310 315 320
 - Ser Ala Arg Val Cys Asp Leu Val Ser Tyr Ile Ala Ala Gln Asp Ala 325 330 335
- 70 Lys Ala

	(2) INFORMATION FOR SEQ ID NO: 11:	
5	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 388 base pairs (B) TYPE: nucleic acid (C) STRANDEINESS: double (D) TOPOLOGY: linear	
10	(iii) MOLECULE TYPE: DNA (genomic)	
	(iii) HYPOTHETICAL: NO	
	(iv) ANTI-SENSE: NO	
15	(vi) ORIGINAL SOURCE: (A) ORGANISM: Phaffia rhodozyma	
20	(ix) FEATURE: (A) NAME/KEY: promoter (B) LOCATION:1385	
25	(ix) FEATURE: (A) NAME/KEY: TATA signal (B) LOCATION:249263 (D) OTHER INFORMATION:/label= putative	
30	<pre>(ix) FEATURE: (A) NAME/KEY: misc_signal (B) LOCATION:287302 (D) OTHER INFORMATION:/function= "cap-signal" /label= putative</pre>	
35	<pre>(ix) FEATURE: (A) NAME/KEY: misc_RNA (B) LOCATION:386388 (D) OTHER INFORMATION:/function= "start of CDS"</pre>	
40	<pre>(ix) FEATURE: (A) NAME/KEY: misc_feature (B) LOCATION:85 (D) OTHER INFORMATION:/note= "uncertain"</pre>	
45	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 11:	
4 5	TOGIGGGIGC ATGIATGIAC GIGAGIGAGI GOOOGGGAAA GGCGAGIACG TGIGIGIACG	60
	CGCAAGGAAG AACAACGAAG CGCANGCTAT GAGCAAGCAC AACTGGGCAC CGAACGAGAA	120
50	CAGIAACIGI COGIATCITIC CCACCGACAC GAGGGGICIC CCGGCGGCAA CCGCCGGIGC	180
	CCCCCICCCC TTACGICAGC CACCCAGITT TCTTCCATCT CTTTCTCTCT CCTTCCAAAA	240
55	GICTTICAGT TITAAACOGC CCCCAAAAAA AGAAGAGGG ACTITITCIT TCCTTCTCTC	300
	CCATCATCCA CAAAGATCTC TCTTCTTCAA CAACAACTAC TACTACTACC ACTACCACCA	360
	CTACTICICT AACACTCTTA CCATCATG	388
60	(2) INFORMATION FOR SEQ ID NO:12:	
65	(i) SEQUENCE CHARACTERISTICS: (A) LENGIH: 2546 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: double (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: cDNA	

		(iii)	HY	POTHE	anc.	AL: 1	3 0										
		(iv)	AN.	ri-si	NSE:	: 100											
5		(vi)	OR:	IGIN A) OF	AL SO RGANT	OURCE	E: Phaf	fia	rho	lozyr	na						
10		(ix)	(<i>I</i>	ATURI A) NV B) L(D) OI	ME/I	CN:	225.	. 224 TON:	16 : /pr	roduc	:t= '	'PRC1	tB"				
		(xi)	SEX	CUENC	E DE	SCRI	PTIC	N: 5	SEQ 1	D 187):12:	:					
15	TCIZ	GAAC	MA (TIGG	ATCCC	x a	eggc1	GCAC	GAZ	ATTCC	GCA	CGAC	CGGZ	AA.	ZAAGZ	AGIGG	60
	ACAC	AGAC	SAG A	ALCLI	MGC	rg az	GAGI	1G12	TTC	CAG	VAAG	GGAZ	AACZ	AA G	GAA	GAAGC	120
20	GCCC	CAAGC	AC A	ATCAC	CAAC	TT	AGC#	AGCC	ල ල	CCAC	3CCC	GATO	TCGC	AT A	GAC	ATCATC	180
20	TIAC	CCAF	CT ()GIAI	CATC	.c a	ZAACZ	AGAT7	A GAC	TTT	ngr	CCC2		Thr		CIC Leu	236
25	GCA Ala 5	TAT Tyr	TAC Tyr	CAG Gln	ATC Ile	CAT His 10	CIG Leu	ATC	TAT Tyr	ACT Thr	CTC Leu 15	CCA Pro	ATT Ile	CTT Leu	GGT Gly	CTT Leu 20	284
30	CTC Leu	Gly	CIG Leu	CTC Leu	ACT Thr 25	TCC Ser	CCG Pro	ATT Ile	TIG Leu	ACA Thr 30	aaa Lys	TTT Phe	GAC Asp	ATC Ile	TAC Tyr 35	AAA Lys	332
35	ATA Ile	TCG Ser	ATC Ile	CTC Leu 40	GTA Val	TTT Phe	ATT Ile	GCG Ala	TTT Phe 45	AGT Ser	GCA Ala	ACC Thr	ACA Thr	CCA Pro 50	TGG Trp	GAC Asp	380
40	TCA Ser	TGG Trp	ATC Ile 55	ATC Ile	AGA Arg	AAT Asn	GGC Gly	GCA Ala 60	TGG Trp	ACA Thr	TAT Tyr	CCA Pro	TCA Ser 65	GCG Ala	GAG Glu	AGT Ser	428
	GJ y GGC	CAA Gln 70	GGC GC	GTG Val	TIT Phe	GGA Gly	ACG Thr 75	TTT Phe	CTA Leu	GAT Asp	GTT Val	CCA Pro 80	TAT Tyr	GAA Glu	GAG Glu	TAC Tyr	476
45	GCT Ala 85	TTC Phe	TTT Phe	GTC Val	ATT Ile	CAA Gln 90	ACC Thr	GTA Val	ATC Ile	ACC Thr	GGC Gly 95	TTG Leu	GTC Val	TAC Tyr	GTC Val	TTG Leu 100	524
50	GCA Ala	ACT Thr	AGG Arg	CAC His	CTT Leu 105	CTC Leu	CCA Pro	TCT Ser	CTC Leu	GCG Ala 110	CTT Leu	CCC Pro	AAG Lys	ACT Thr	AGA Arg 115	TCG Ser	5 7 2
55	TCC Ser	GCC Ala	CTT Leu	TCT Ser 120	CTC Leu	GOG Ala	CTC Leu	AAG Lys	GOG Ala 125	CIC Leu	ATC Ile	CCT Pro	CTG Leu	CCC Pro 130	ATT Ile	ATC Ile	620
60	TAC Tyr	CTA Leu	TTT Phe 135	ACC Thr	GCT Ala	CAC His	CCC Pro	AGC Ser 140	CCA Pro	TCG Ser	CCC Pro	gac Asp	CCG Pro 145	CIC Leu	GTG Val	ACA Thr	66 8
•	GAT Asp	CAC His 150	TAC Tyr	TTC Phe	TAC Tyr	ATG Met	CGG Arg 155	GCA Ala	CTC Leu	TCC Ser	TTA Leu	CTC Leu 160	ATC Ile	ACC Thr	CCA Pro	CCT Pro	716
65	ACC Thr 165	ATG Met	CTC Leu	TTG Leu	GCA Ala	GCA Ala 170	TTA Leu	TCA Ser	GGC Gly	GAA Glu	TAT Tyr 175	GCT Ala	TTC Phe	GAT Asp	TGG Trp	AAA Lys 180	764
70	AGT Ser	GGC Gly	CGA Arg	GCA Ala	AAG Lys	TCA Ser	ACT Thr	ATT Ile	GCA Ala	GCA Ala	ATC Ile	ATG Met	ATC Ile	CCG Pro	ACG Thr	GIG Val	812

	WC	97/:	2363:	3								^					PCT
											4	9					
					185					190					195		
5	TAT Tyr	CIG	ATT	TGG Trp 200	Val	GAT Asp	TAT	GIT Val	GCT Ala 205	Val	Gly	CAA Gln	GAC Asp	Ser 210	Trp	TCG Ser	860
10	ATC	AAC Asn	GAT Asp 215	GAG Glu	AAG Lys	ATT	GTA Val	GGG Gly 220	TGG Trp	AGG Arg	CTT	GGA Gly	GGT Gly 225	Val	CTA Leu	CCC Pro	908
	ATT Ile	GAG Glu 230	GAA Glu	GCT Ala	ATG Met	TTC Phe	TTC Phe 235	TTA Leu	CTG Leu	ACG Thr	AAT Asn	CIA Leu 240	ATG Met	ATT	GTT Val	CTG Leu	956
15	GGT Gly 245	CIG Leu	TCT Ser	GCC Ala	TGC Cys	GAT Asp 250	CAT His	ACT Thr	CAG Gln	GCC Ala	CTA Leu 255	TAC Tyr	CTG Leu	CIA Leu	CAC His	GGT Gly 260	1004
20	CGA Arg	ACT Thr	ATT	TAT Tyr	GGC Gly 265	AAC Asn	AAA Lys	AAG Lys	ATG Met	CCA Pro 270	TCT Ser	TCA Ser	TTT Phe	CCC Pro	CIC Leu 275	ATT Ile	1052
25	ACA Thr	CCG Pro	CCT Pro	GTG Val 280	CIC Leu	TCC Ser	CIG Leu	TTT Phe	TTT Phe 285	AGC Ser	AGC Ser	CGA Arg	CCA Pro	TAC Tyr 290	TCT Ser	TCT Ser	1100
30	Gln CAG	CCA Pro	AAA Lys 295	CGT Arg	GAC Asp	TIG Leu	GAA Glu	CTG Leu 300	GCA Ala	GTC Val	AAG Lys	TTG Leu	TTG Leu 305	GAG Glu	AAA Lys	AAG Lys	1148
	AGC Ser	CGG Arg 310	AGC Ser	TTT Phe	TTT Phe	GTT Val	GCC Ala 315	TCG Ser	GCT Ala	GGA Gly	TTT Phe	CCT Pro 320	AGC Ser	GAA Glu	GTT Val	AGG Arg	1196
35	GAG Glu 325	AGG Arg	CTG Leu	GTT Val	GGA Gly	CIA Leu 330	TAC Tyr	GCA Ala	TTC Phe	TGC Cys	CGG Arg 335	GTG Val	ACT Thr	GAT Asp	GAT Asp	CTT Leu 340	1244
40	ATC Ile	GAC Asp	TCT Ser	CCT Pro	GAA Glu 345	GTA Val	TCT Ser	TCC Ser	AAC Asn	CCG Pro 350	CAT His	GCC Ala	ACA Thr	ATT Ile	GAC Asp 355	ATG Met	1292
45	GTC Val	TCC Ser	GAT Asp	TTT Phe 360	CTT Leu	ACC Thr	CTA Leu	CTA Leu	TTT Phe 365	GGG Gly	CCC Pro	CCG Pro	CTA Leu	CAC His 370	CCT Pro	TCG Ser	1340
50	CAA Gln	CCT Pro	GAC Asp 375	AAG Lys	ATC Ile	CTT Leu	TCT Ser	TCG Ser 380	CCT Pro	TTA Leu	CTT Leu	CCT Pro	CCT Pro 385	TCG Ser	CAC His	CCT Pro	1388
55	TCC Ser	CGA Arg 390	Pro	ACG Thr	GGA Gly	ATG Met	TAT Tyr 395	CCC Pro	CTC Leu	CCG Pro	CCT Pro	CCT Pro 400	CCT Pro	TCG Ser	CIC Leu	TCG Ser	1436
	CCT Pro 405	GCC Ala	GAG Glu	CTC Leu	GTT Val	CAA Gln 410	TTC Phe	CTT Leu	ACC Thr	GAA Glu	AGG Arg 415	GTT Val	CCC Pro	GTT Val	CAA Gln	TAC Tyr 420	1484
60	CAT His	TTC Phe	GCC Ala	TTC Phe	AGG Arg 425	TTG Leu	CIC Leu	GCT Ala	AAG Lys	TIG Leu 430	CAA Gln	GGG Gly	CIG Leu	ATC Ile	CCT Pro 435	CGA Arg	1532
65	TAC Tyr	CCA Pro	CTC Leu	GAC Asp 440	GAA Glu	CTC Leu	CTT Leu	AGA Arg	GGA Gly 445	TAC Tyr	ACC Thr	ACT Thr	GAT Asp	CTT Leu 450	ATC Ile	TTT Phe	1580
	~~~																

# SUBSTITUTE SHEET (RULE 26)

1628

CCC TTA TCG ACA GAG GCA GTC CAG GCT CGG AAG ACG CCT ATC GAG ACC Pro Leu Ser Thr Glu Ala Val Gln Ala Arg Lys Thr Pro Ile Glu Thr 455 460 465

	ACA Thr																1676
5	GAG Glu 485			GIC Val													1724
10				GAA Glu													1772
15				CTT Leu 520												yab GyC	1820
20				GGG Gly													1868
20	GAT Asp	GAA Glu 550	Ser	AAG Lys	CTT Leu	GCG Ala	ATC Ile 555	CCG Pro	ACT Thr	GAT Asp	TCG Trp	ACG Thr 560	Glu	CCT Pro	CGG Arg	CCT Pro	1916
25		Asp		GAC Asp			Leu					Ser				CCA Pro 580	1964
30				GCC Ala		Glu											2012
35	CTI Leu	Pro	TTA	GTC Val 600	Ala	TAC	GCA Ala	GAG Glu	GAT Asp 605	Leu	GCC	AAA Lys	CAI His	TCI Ser 610	Tyx	'AAG 'Lys	2060
40	GCP	ATT	GAC Asp 619	) Arg	CTI Leu	CCI Pro	ACC Thr	GAG Glu 620	Val	CAA Glr	GCC Ala	GCZ Gly	A ATG / Met 625	Arg	GCG Ala	GCT Ala	2108
45	TGC Cys	GCC 630	a Sei	TAC Tyr	CIA Lev	CIC Let	ATC 11e 635	e Gly	CGA Arg	GAC Glu	ATC	E AA E Ly: 640	s Val	C GM L Val	TIGO L Trip	AAA D Lys	2156
	Gly	A GAN Y Asi 5	p Va	l Gly	y Gli	ı Arş	A ACK g Arg 0	y Tîn	r Val	L Ala	a Gl	y Tr	p Arg	J Ary	y Val	A CGG 1 Arg 660	2204
50	AA: Ly:	A GT s Va	C TN l Le	GAG uSe:	r GN r Va 66	l Va	CAT 1 Me	G AGK	c GG c Gl	A TG y Trj 67	o Gl	A GG u Gl	G CA y Gl	g TA	AGAC	AGCG	2253
55																TICTCIA	2313
																GATATTT	2373
60																TTAGAAT	2493
	ניד	ATA	TTA	ACZ	AGGT1	ATA	GACC	ATA	AG A	CTAP	IAAA	K A	<b>LAAA</b>	AAA	AA A		2546

## (2) INFORMATION FOR SEQ ID NO:13:

## (i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 673 amino acids
(B) TYPE: amino acid

(D) TOPOLOGY: linear

- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:13:
- Met Thr Ala Leu Ala Tyr Tyr Gln Ile His Leu Ile Tyr Thr Leu Pro 1 5 10 15
  - Ile Leu Gly Leu Leu Gly Leu Leu Thr Ser Pro Ile Leu Thr Lys Phe
    20 25 30
- Asp Ile Tyr Lys Ile Ser Ile Leu Val Phe Ile Ala Phe Ser Ala Thr 35 40 45
- Thr Pro Trp Asp Ser Trp Ile Ile Arg Asn Gly Ala Trp Thr Tyr Pro
  - Ser Ala Glu Ser Gly Gln Gly Val Phe Gly Thr Phe Leu Asp Val Pro 65 70 75 80
- Tyr Glu Glu Tyr Ala Phe Phe Val Ile Gln Thr Val Ile Thr Gly Leu 85 90 95
  - Val 7,7 Val Leu Ala Thr Arg His Leu Leu Pro Ser Leu Ala Leu Pro 100 105 110
  - Lys Thr Arg Ser Ser Ala Leu Ser Leu Ala Leu Lys Ala Leu Ile Pro 115 120 125
- Lew Pro Ile Ile Tyr Lew Phe Thr Ala His Pro Ser Pro Ser Pro Asp 130 135 140
  - Pro Leu Val Thr Asp His Tyr Phe Tyr Met Arg Ala Leu Ser Leu Leu 145 150 155 160
- Ile Thr Pro Pro Thr Met Leu Leu Ala Ala Leu Ser Gly Glu Tyr Ala 165 170 175
  - Phe Asp Trp Lys Ser Gly Arg Ala Lys Ser Thr Ile Ala Ala Ile Met 180 185 190
- Ile Pro Thr Val Tyr Leu Ile Trp Val Asp Tyr Val Ala Val Gly Gln
  195 200 205
- Asp Ser Trp Ser Ile Asm Asp Glu Lys Ile Val Gly Trp Arg Leu Gly
  210 215 220
  - Gly Val Leu Pro Ile Glu Glu Ala Met Phe Phe Leu Leu Thr Asn Leu 225 230 235 240
- Met Ile Val Leu Gly Leu Ser Ala Cys Asp His Thr Gln Ala Leu Tyr 245 250 255
  - Leu Leu His Gly Arg Thr Ile Tyr Gly Asn Lys Lys Met Pro Ser Ser 260 265 270
- Phe Pro Leu Ile Thr Pro Pro Val Leu Ser Leu Phe Phe Ser Ser Arg 275 280 285
- Pro Tyr Ser Ser Gln Pro Lys Arg Asp Leu Glu Leu Ala Val Lys Leu 290 295 300
  - Leu Glu Lys Lys Ser Arg Ser Phe Phe Val Ala Ser Ala Gly Phe Pro 305 310 315 320
- 65 Ser Glu Val Arg Glu Arg Leu Val Gly Leu Tyr Ala Phe Cys Arg Val 325 330 335
  - Thr Asp Asp Leu Ile Asp Ser Pro Glu Val Ser Ser Asn Pro His Ala 340 345 350

Thr Ile	Asp	Met	Val	Ser	Asp	Phe	Leu	Thr	Leu	Leu	Phe	Gly	Pro	Pro
	355					360					365			

- Leu His Pro Ser Gln Pro Asp Lys Ile Leu Ser Ser Pro Leu Leu Pro 370 375 380
  - Pro Ser His Pro Ser Arg Pro Thr Gly Met Tyr Pro Leu Pro Pro 385 390 395 400
- Pro Ser Leu Ser Pro Ala Glu Leu Val Gln Phe Leu Thr Glu Arg Val 405 410 415
  - Pro Val Gln Tyr His Phe Ala Phe Arg Leu Leu Ala Lys Leu Gln Gly 420 425 430
  - Leu Ile Pro Arg Tyr Pro Leu Asp Glu Leu Leu Arg Gly Tyr Thr Thr 435 440 445
- Asp Leu Ile Phe Pro Leu Ser Thr Glu Ala Val Gln Ala Arg Lys Thr 20 450 455 460
  - Pro Ile Glu Thr Thr Ala Asp Leu Leu Asp Tyr Gly Leu Cys Val Ala 465 470 475 480
- Gly Ser Val Ala Glu Leu Leu Val Tyr Val Ser Trp Ala Ser Ala Pro 485 490 495
  - Ser Gln Val Pro Ala Thr Ile Glu Glu Arg Glu Ala Val Leu Val Ala 500 505 510
- Ser Arg Glu Met Gly Thr Ala Leu Gln Leu Val Asn Ile Ala Arg Asp 515 520 525
- Ile Lys Gly Asp Ala Thr Glu Gly Arg Phe Tyr Leu Pro Leu Ser Phe 535 540
  - Phe Gly Leu Arg Asp Glu Ser Lys Leu Ala Ile Pro Thr Asp Trp Thr 545 550 555 560
- 40 Glu Pro Arg Pro Gln Asp Phe Asp Lys Leu Leu Ser Leu Ser Pro Ser 565 570 575
  - Ser Thr Leu Pro Ser Ser Asn Ala Ser Glu Ser Phe Arg Phe Glu Trp
    580 585 590
  - Lys Thr Tyr Ser Leu Pro Leu Val Ala Tyr Ala Glu Asp Leu Ala Lys 595 600 605
- His Ser Tyr Lys Gly Ile Asp Arg Leu Pro Thr Glu Val Gln Ala Gly 610 615 620
  - Met Arg Ala Ala Cys Ala Ser Tyr Leu Leu Ile Gly Arg Glu Ile Lys 625 630 635 640
- 55 Val Val Trp Lys Gly Asp Val Gly Glu Arg Arg Thr Val Ala Gly Trp 645 650 655
  - Arg Arg Val Arg Lys Val Leu Ser Val Val Met Ser Gly Trp Glu Gly 660 665 670

Gln

60

65

45

#### (2) INFORMATION FOR SEQ ID NO:14:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGIH: 1882 base pairs
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: double
- (D) TOPOLOGY: linear

	(ii) MOLECULE TYP	PE: CDNA	
	(iii) HYPOTHETICAL	: NO	
5	(iv) ANTI-SENSE: 1	NO	
	(vi) ORIGINAL SOU (A) ORGANIS	RCE: M: Phaffia rhodozyma	
10	(ix) FEATURE: (A) NAME/KE: (B) LOCATION (D) OTHER IN		
15	(xi) SEQUENCE DESC	CRIPTION: SEQ ID NO:14:	
	GGCACGAGCC AATTIAAAGT	GCACTCAGCC ATAGCTAACA CACAGAACTA CACATACATA	60
20	CACTCATCOG GAACACATAG	G ATG GAT TAC GCG AAC ATC CTC ACA GCA ATT Met Asp Tyr Ala Asm Ile Leu Thr Ala Ile 1 5 10	111
25	CCA CTC GAG TIT ACT CO Pro Leu Glu Phe Thr Pi 15	CT CAG GAT GAT ATC GTG CTC CTT GAA CCG TAT TO Gln Asp Asp Ile Val Leu Leu Glu Pro Tyr 20 25	159
70	CAC TAC CTA GGA AAG AA His Tyr Leu Gly Lys As 30	AC CCT GGA AAA GAA ATT CGA TCA CAA CTC ATC sn Pro Gly Lys Glu Ile Arg Ser Gln Leu Ile 35 40	207
30	GAG GCT TTC AAC TAT TO Glu Ala Phe Asn Tyr To 45	GG TTG GAT GTC AAG AAG GAG GAT CTC GAG GTC TP Leu Asp Val Lys Lys Glu Asp Leu Glu Val 50 55	255
35	ATC CAG AAC GIT GIT G Ile Gln Asn Val Val G 60	GC ATG CTA CAT ACC GCT AGC TTA TTA ATG GAC ly Met Leu His Thr Ala Ser Leu Leu Met Asp 65 70	303
40	Asp Val Glu Asp Ser Se	CG GTC CTC AGG CGT GGG TCG CCT GTG GCC CAT er Val Leu Arg Arg Gly Ser Pro Val Ala His 80 85 90	351
45	CTA ATT TAC GGG ATT CO Leu Ile Tyr Gly Ile P: 95	TCG CAG ACA ATA AAC ACT GCA AAC TAC GTC TAC TO Gln Thr Ile Asn Thr Ala Asn Tyr Val Tyr 100 105	399
50	TIT CIG GCT TAT CAA G Phe Leu Ala Tyr Gln G 110	AG ATC TTC AAG CTT CGC CCA ACA CCG ATA CCC Flu Ile Phe Lys Leu Arg Pro Thr Pro Ile Pro 115 120	447
20	ATG CCT GTA ATT CCT C Met Pro Val Ile Pro P 125	CT TCA TCT GCT TCG CTT CAA TCA TCC GTC TCC Pro Ser Ser Ala Ser Leu Gln Ser Ser Val Ser 130 135	495
55	TCT GCA TCC TCC TCC T Ser Ala Ser Ser Ser Se 140	CC TCG GCC TCG TCT GAA AAC GGG GGC ACG TCA Ser Ser Ala Ser Ser Glu Asm Gly Gly Thr Ser 145	543
60	Thr Pro Asn Ser Gln I	ATT CCG TTC TCG AAA GAT ACG TAT CTT GAT AAA Cle Pro Phe Ser Lys Asp Thr Tyr Leu Asp Lys 165 170	591
65	GTG ATC ACA GAC GAG A Val Ile Thr Asp Glu M 175	ATG CTT TCC CTC CAT AGA GGG CAA GGC CTG GAG Met Leu Ser Leu His Arg Gly Gln Gly Leu Glu 180 185	639
	CTA TTC TGG AGA GAT A Leu Phe Trp Arg Asp S 190	AGT CTG ACG TGT CCT AGC GAA GAG GAA TAT GTG Ser Leu Thr Cys Pro Ser Glu Glu Glu Tyr Val 195 200	687

(2) INFORMATI	ON FOR	SEQ	$\mathbf{I}$	NO:15:
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## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGIH: 376 amino acids
- (B) TYPE: amino acid
- (D) TOPOLOGY: linear

#### (ii) MOLECULE TYPE: protein

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:15:

Met Asp Tyr Ala Asn Ile Leu Thr Ala Ile Pro Leu Glu Phe Thr Pro 1 5 10 15

Gln Asp Asp Ile Val Leu Leu Glu Pro Tyr His Tyr Leu Gly Lys Asn 20 25 30

Pro Gly Lys Glu Ile Arg Ser Gln Leu Ile Glu Ala Phe Asn Tyr Trp 35 40 45

Leu Asp Val Lys Lys Glu Asp Leu Glu Val Ile Gln Asn Val Val Gly 50 55 60

Met Leu His Thr Ala Ser Leu Leu Met Asp Asp Val Glu Asp Ser Ser 5 65 70 75 80

Val Leu Arg Arg Gly Ser Pro Val Ala His Leu Ile Tyr Gly Ile Pro 85 90 95

30 Gln Thr Ile Asn Thr Ala Asn Tyr Val Tyr Phe Leu Ala Tyr Gln Glu 100 105 110

Ile Phe Lys Leu Arg Pro Thr Pro Ile Pro Met Pro Val Ile Pro Pro 115 120 125

Ser Ser Ala Ser Leu Gln Ser Ser Val Ser Ser Ala Ser Ser Ser Ser 130 140

40 Ser Ala Ser Ser Glu Asn Gly Gly Thr Ser Thr Pro Asn Ser Gln Ile 145 150 155 160

Pro Phe Ser Lys Asp Thr Tyr Leu Asp Lys Val Ile Thr Asp Glu Met 165 170 175

Leu Ser Leu His Arg Gly Gln Gly Leu Glu Leu Phe Trp Arg Asp Ser 180 185 190

Leu Thr Cys Pro Ser Glu Glu Glu Tyr Val Lys Met Val Leu Gly Lys 195 200 205

Thr Gly Gly Leu Phe Arg Ile Ala Val Arg Leu Met Met Ala Lys Ser 210 215 220

Glu Cys Asp Ile Asp Phe Val Gln Leu Val Asn Leu Ile Ser Ile Tyr 225 230 235 240

Phe Gln Ile Arg Asp Asp Tyr Met Asn Leu Gln Ser Ser Glu Tyr Ala 245 250 255

His Asn Lys Asn Phe Ala Glu Asp Leu Thr Glu Gly Lys Phe Ser Phe 260 265 270

Pro Thr Ile His Ser Ile His Ala Asn Pro Ser Ser Arg Leu Val Ile 275 280 285

Asn Thr Leu Gln Lys Lys Ser Thr Ser Pro Glu Ile Leu His His Cys 290 295 300

 ${\mathfrak w}$  Val Asn Tyr Met Arg Thr Glu Thr His Ser Phe Glu Tyr Thr Gln Glu

	305					310					315					32	20		
5	Val 1	Leu .	Asn	Thr	Leu 325	Ser	Gly	Ala		Glu 330	Arg	Glu	Leu	Gly	Arg 335		eu		
,	Gln (	Gly	Glu	Phe 340	Ala	Glu .	Ala		Ser 345	Arg	Met	Asp	Leu	Gly 350	_	Vā	al		
0	Asp :		Glu 355	Gly	Arg	Thr		Lys . 360	Asn	Val	Lys	Leu	Glu 365	Ala	Ile	L	eu		
	Lys :	Lys 370	Leu	Ala	Asp	Ile	Pro 375	Leu											
5	(2)	INFC	RMAT	.IOM	FOR	SEQ	m n	10:16	:										
NO		(i)	(E (C	() LE () T'( () S'1	NGIH PE:	i: 20 nucl EDNE	58 b eic SS:	STIC ase acid doub ar	pair [	s									
ಶ	(				E TY			<b>\</b>											
10			ORI	GIN	NSE:	OURCE		ffia	*		••								
35		(ix)	FE/ () (I	ATURI A) N B) L	E: AME/I CCATI	ŒY: ION:	CDS 46.	. 1794 PION :	l į			"PRC	rtI"						
40	CCTC							ON: S	_						KGA A				54
45	GAA Glu	CAA Gln 5	Asp	CAG Gln	GAT Asp	AAA Lys	CCC Pro 10	Thr	GCT Ala	ATC	ATC	GIG Val	Gl	TG	r GG.	r P	ATC .	1	L02
50							Ala	CGT Arg				Glu						;	150
55						Asn		TAC Tyr			Gly				r Le			:	198
	_			_	/ Tyr		_	GAT Asp	_	Gly				ı Le				:	246
60				ı Phe				TTC Phe	Glu					u Ly					294
65			va.					ing Cys				а Ту							342
	CAC	GA	r GA	A GA	G AC		C AC	r TT	r TO	A AC		CAI	c cc	G T	rs cr	rc	AAG		390

	100					105					110					115	
5	COG Arg	GAA Glu	GTC Val	GAG Glu	CGT Arg 120	TTT Phe	GAA Glu	GC Gly	AAA Lys	GAT Asp 125	GGA Gly	TTT Phe	GAT Asp	CGG Arg	TTC Phe 130	TTG Leu	438
10	TCG Ser	TTT Phe	ATC Ile	CAA Gln 135	GAA Glu	GCC Ala	CAC His	AGA Arg	CAT His 140	TAC Tyr	GAG Glu	CTT Leu	GCT Ala	GTC Val 145	GIT Val	CAC His	486
10	GTC Val	CTG Leu	CAG Gln 150	AAG Lys	AAC Asn	TTC Phe	CCT Pro	GGC Gly 155	TTC Phe	GCA Ala	GCA Ala	TTC Phe	TTA Leu 160	CGG Arg	CTA Leu	CAG Gln	534
15	TTC Phe	ATT Ile 165	GGC Gly	CAA Gln	ATC Ile	CTG Leu	GCT Ala 170	CTT Leu	CAC His	CCC Pro	TTC Phe	GAG Glu 175	TCT Ser	ATC Ile	TGG Trp	ACA Thr	582
20	AGA Arg 180	GTT Val	IGT Cys	CGA Arg	TAT Tyr	TTC Phe 185	aag Lys	ACC Thr	GAC Asp	AGA Arg	TTA Leu 190	CGA Arg	AGA Arg	GTC Val	TTC Phe	TCG Ser 195	630
25	TTT Phe	GCA Ala	GTG Val	ATG Met	TAC Tyr 200	ATG Met	GGT Gly	CAA Gln	AGC Ser	CCA Pro 205	TAC Tyr	AGT Ser	GCG Ala	CCC Pro	GGA Gly 210	ACA Thr	678
30	TAT Tyr	TCC Ser	TTG Leu	CTC Leu 215	CAA Gln	TAC Tyr	ACC Thr	GAA Glu	TTG Leu 220	ACC Thr	GAG Glu	GC Gly	ATC Ile	TGG Trp 225	TAT Tyr	CCG Pro	726
	AGA Arg	GGA Gly	GGC Gly 230	TTT Phe	TGG Trp	CAG Gln	GTT Val	CCT Pro 235	AAT Asn	ACT Thr	CIT Leu	CTT Leu	CAG Gln 240	ATC Ile	GTC Val	AAG Lys	774
35	CGC Arg	AAC Asn 245	AAT Asn	CCC Pro	TCA Ser	GCC Ala	AAG Lys 250	TIC Phe	AAT Asn	TIC Phe	AAC Asn	GCT Ala 255	CCA Pro	GTT Val	TCC Ser	CAG Gln	822
40	GIT Val 260	CTT Leu	CIC Leu	TCT Ser	CCT Pro	GCC Ala 265	AAG Lys	GAC Asp	CGA Arg	GCG Ala	ACT Thr 270	GT Gly	GTT Val	CGA Arg	CTT Leu	GAA Glu 275	870
45	TCC Ser	GGC Gly	GAG Glu	GAA Glu	CAT His 280	CAC His	GCC Ala	GAT Asp	GIT Val	GTG Val 285	ATT Ile	GIC Val	AAT Asn	GCT Ala	GAC Asp 290	CTC Leu	918
50	Val	Тух	Ala	Ser 295	Glu	His	Leu	Ile	Pro 300	Asp	GAT Asp	Ala	Arg	Asn 305	Lys	Ile	966
	Gly	Gln	Leu 310	Gly	Glu	Val	Lys	Arg 315	Ser	Trp	TGG Trp	Ala	320	Leu	Val	Gly	1014
55	GGA Gly	AAG Lys 325	AAG Lys	CIC Leu	AAG Lys	GGA Gly	AGT Ser 330	TGC Cys	AGT Ser	AGT Ser	TIG Leu	AGC Ser 335	TTC Phe	TAC Tyr	TGG Trp	AGC Ser	1062
60	ATG Met 340	Asp	CGA Arg	ATC Ile	GIG Val	GAC Asp 345	ggr Gly	CIG Leu	GCC	GGA Gly	CAC His 350	AAT Asn	ATC	TTC Phe	TTG Leu	GCC Ala 355	1110
65	GAG Glu	GAC Asp	TTC Phe	AAG Lys	GGA Gly 360	TCA Ser	TTC Phe	GAC Asp	ACA Thr	ATC Ile 365	TTC Phe	GAG Glu	GAG Glu	TIG	GGT Gly 370	CTC Leu	1158
70	CCA Pro	GCC Ala	GAT Asp	CCT Pro 375	TCC Ser	TTT Phe	TAC Tyr	GTG Val	AAC Asn 380	GTT Val	Pro	TCG Ser	CGA Arg	ATC Ile 385	Asp	CCT Pro	1206

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO:17:

Met Gly Lys Glu Gln Asp Gln Asp Lys Pro Thr Ala Ile Ile Val Gly Cys Gly Ile Gly Gly Ile Ala Thr Ala Ala Arg Leu Ala Lys Glu Gly Phe Gln Val Thr Val Phe Glu Lys Asn Asp Tyr Ser Gly Gly Arg Cys Ser Leu Ile Glu Arg Asp Gly Tyr Arg Phe Asp Gln Gly Pro Ser Leu 55 Leu Leu Pro Asp Leu Phe Lys Gln Thr Phe Glu Asp Leu Gly Glu Lys Met Glu Asp Trp Val Asp Leu Ile Lys Cys Glu Pro Asn Tyr Val Cys His Phe His Asp Glu Glu Thr Phe Thr Phe Ser Thr Asp Met Ala 105 Leu Leu Lys Arg Glu Val Glu Arg Phe Glu Gly Lys Asp Gly Phe Asp Arg Phe Leu Ser Phe Ile Gln Glu Ala His Arg His Tyr Glu Leu Ala Val Val His Val Leu Gln Lys Asn Phe Pro Gly Phe Ala Ala Phe Leu Arg Leu Gln Phe Ile Gly Gln Ile Leu Ala Leu His Pro Phe Glu Ser Ile Trp Thr Arg Val Cys Arg Tyr Phe Lys Thr Asp Arg Leu Arg Arg Val Phe Ser Phe Ala Val Met Tyr Met Gly Gln Ser Pro Tyr Ser Ala Pro Gly Thr Tyr Ser Leu Leu Gln Tyr Thr Glu Leu Thr Glu Gly Ile 215 Trp Tyr Pro Arg Gly Gly Phe Trp Gln Val Pro Asn Thr Leu Leu Gln Ile Val Lys Arg Asn Asn Pro Ser Ala Lys Phe Asn Phe Asn Ala Pro Val Ser Gln Val Leu Leu Ser Pro Ala Lys Asp Arg Ala Thr Gly Val Arg Leu Glu Ser Gly Glu Glu His His Ala Asp Val Val Ile Val Asn 280 Ala Asp Leu Val Tyr Ala Ser Glu His Leu Ile Pro Asp Asp Ala Arg Asn Lys Ile Gly Gln Leu Gly Glu Val Lys Arg Ser Trp Trp Ala Asp Leu Val Gly Gly Lys Lys Leu Lys Gly Ser Cys Ser Ser Leu Ser Phe 330 Tyr Trp Ser Met Asp Arg Ile Val Asp Gly Leu Gly Gly His Asn Ile Phe Leu Ala Glu Asp Phe Lys Gly Ser Phe Asp Thr Ile Phe Glu Glu

Leu Gly Leu Pro Ala Asp Pro Ser Phe Tyr Val Asn Val Pro Ser Arg Ile Asp Pro Ser Ala Ala Pro Glu Gly Lys Asp Ala Ile Val Ile Leu 390 Val Pro Cys Gly His Ile Asp Ala Ser Asn Pro Gln Asp Tyr Asn Lys Leu Val Ala Arg Ala Arg Lys Phe Val Ile Gln Thr Leu Ser Ala Lys Leu Gly Leu Pro Asp Phe Glu Lys Met Ile Val Ala Glu Lys Val His 440 15 Asp Ala Pro Ser Trp Glu Lys Glu Phe Asn Leu Lys Asp Gly Ser Ile Leu Gly Leu Ala His Asn Phe Met Gln Val Leu Gly Phe Arg Pro Ser 20 Thr Arg His Pro Lys Tyr Asp Lys Leu Phe Phe Val Gly Ala Ser Thr 485 His Pro Gly Thr Gly Val Pro Ile Val Leu Ala Gly Ala Lys Leu Thr Ala Asn Gln Val Leu Glu Ser Phe Asp Arg Ser Pro Ala Pro Asp Pro 520 30 Asn Met Ser Leu Ser Val Pro Tyr Gly Lys Pro Leu Lys Ser Asn Gly Thr Gly Ile Asp Ser Gln Val Gln Leu Lys Phe Met Asp Leu Glu Arg 35 Trp Val Tyr Leu Leu Val Leu Leu Ile Gly Ala Val Ile Ala Arg Ser 565

45 (2) INFORMATION FOR SEQ ID NO:18:

Val Gly Val Leu Ala Phe 580

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGIH: 2470 base pairs
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: double
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: cDNA
- 55 (iii) HYPOTHETICAL: NO
  - (iv) ANTI-SENSE: NO
  - (vi) ORIGINAL SOURCE:
  - (A) ORGANISM: Phaffia rhodozyma
    - (ix) FEATURE:
      - (A) NAME/KEY: CDS
      - (B) LOCATION: 177..2198
      - (D) OTHER INFORMATION: /product= "PRCITY"
    - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:18:

AACAAGAAGI GGACACAGAG AGATCITIGC TGAAGAGITG TATTCCAGAA AGGGAAAACA

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n	
·	4

	AAG	GAAA	GAA (	3CGC	CGAA	GC A	CATC	ACCA	A CT	TCAG	CAAG	ccca	GTCC	AGC (	CCGA'	rcross	120
	ATA	GACA'	TCA '	TCTT	ACCC	AA C	ICGI	ATCA	rca	CCAA	CAGA	TAG	AGTT	TT (	GTCG(	ZA.	176
5	ATG Met 1	ACG Thr	GCT Ala	CTC Leu	GCA Ala 5	TAT Tyr	TAC Tyr	CAG Gln	ATC Ile	CAT His 10	CIG Leu	ATC Ile	TAT Tyr	ACT Thr	CIC Leu 15	CCA Pro	224
10	ATT	CIT Leu	Gly	CTT Leu 20	CTC Leu	GGC Gly	CIG Leu	CIC Leu	ACT Thr 25	TCC Ser	CCG Pro	ATT Ile	TTG Leu	ACA Thr 30	AAA Lys	TTT Phe	272
15	GAC Asp	ATC	TAC Tyr 35	AAA Lys	ATA Ile	TCG Ser	ATC Ile	CIC Leu 40	GTA Val	TTT Phe	ATT Ile	GCG Ala	TTT Phe 45	AGT Ser	GCA Ala	ACC Thr	320
20	ACA Thr	CCA Pro 50	drL	GAC Asp	TCA Ser	TGG Trp	ATC Ile 55	ATC Ile	AGA Arg	AAT Asn	GGC Gly	GCA Ala 60	TGG Trp	ACA Thr	TAT Tyr	CCA Pro	368
	TCA Ser 65	GCG Ala	GAG Glu	AGT Ser	GGC Gly	CAA Gln 70	GGC Gly	GTG Val	TIT Phe	GGA Gly	ACG Thr 75	TTT Phe	CTA Leu	Asp GAT	GTT Val	CCA Pro 80	416
25	TAT Tyr	GAA Glu	GAG Glu	TAC Tyr	GCT Ala 85	TTC Phe	TTT Phe	GTC Val	ATT Ile	CAA Gln 90	ACC Thr	GTA Val	ATC Ile	ACC Thr	GGC Gly 95	TIG Leu	464
30	GTC Val	TAC Tyr	GTC Val	TTG Leu 100	GCA Ala	ACT Thr	AGG Arg	CAC His	CTT Leu 105	CIC Leu	CCA Pro	TCT Ser	CTC Leu	GCG Ala 110	CIT Leu	CCC Pro	512
35	AAG Lys	ACT Thr	AGA Arg 115	TCG Ser	TCC Ser	GCC Ala	CTT Leu	TCT Ser 120	CTC Leu	GCG Ala	CIC Leu	AAG Lys	GCG Ala 125	CIC Leu	ATC Ile	CCT Pro	560
40	CTG Leu	CCC Pro 130	ATT Ile	ATC Ile	TAC Tyr	CIA Leu	TTT Phe 135	ACC Thr	GCT Ala	CAC His	CCC Pro	AGC Ser 140	CCA Pro	TCG Ser	CCC Pro	GAC Asp	608
	CCG Pro 145	CIC Leu	GTG Val	ACA Thr	GAT Asp	CAC His 150	TAC Tyr	TTC Phe	TAC Tyr	ATG Met	CGG Arg 155	GCA Ala	CIC Leu	TCC Ser	TIA Leu	CIC Leu 160	656
45	ATC Ile	ACC Thr	CCA Pro	CCT Pro	ACC Thr 165	ATG Met	CTC Leu	TIG Leu	GCA Ala	GCA Ala 170	TTA Leu	TCA Ser	GGC Gly	GAA Glu	TAT Tyr 175	GCT Ala	704
50	TTC Phe	gat Asp	TGG Trp	AAA Lys 180	AGT Ser	GGC Gly	CGA Arg	GCA Ala	AAG Lys 185	TCA Ser	ACT Thr	ATT Ile	GCA Ala	GCA Ala 190	ATC Ile	ATG Met	752
55	ATC	CCG Pro	ACG Thr 195	GIG Val	TAT Tyr	CIG Leu	ATT Ile	TGG Trp 200	GTA Val	gat Asp	TAT Tyr	GIT Val	GCT Ala 205	GTC Val	GT Gly	CAA Gln	800
60	GAC Asp	TCT Ser 210	TGG Trp	TCG Ser	ATC Ile	AAC Asn	GAT Asp 215	GAG Glu	AAG Lys	ATT Ile	GIA Val	GGG Gly 220	TGG Trp	AGG Arg	CIT Leu	GCA Gly	848
	GGT Gly 225	GTA Val	CTA Leu	CCC Pro	ATT Ile	GAG Glu 230	GAA Glu	GCT Ala	ATG Met	TTC Phe	TIC Phe 235	TIA Leu	CIG Leu	ACG Thr	TAA Asn	CTA Leu 240	896
65	ATG Met	ATT Ile	GIT Val	CTG Leu	GGT Gly 245	CIG Leu	TCT Ser	GCC Ala	TGC Cys	GAT Asp 250	CAT His	ACT Thr	CAG Gln	GCC Ala	CIA Leu 255	TAC Tyr	944
70	CTG Leu	CTA Leu	CAC His	GGT Gly	CGA Arg	ACT Thr	ATT Ile	TAT Tyr	GGC Gly	AAC Asn	aaa Lys	AAG Lys	ATG Met	CCA Pro	TCT Ser	TCA Ser	992

				260					265					270				
5	TTT Phe	CCC Pro	CIC Leu 275	ATT Ile	ACA Thr	CCG Pro	CCT Pro	GTG Val 280	CIC Leu	TCC Ser	CIG Leu	TTT Phe	TTT Phe 285	AGC Ser	AGC Ser	CGA Arg	10	040
10	CCA Pro	TAC Tyr 290	TCT Ser	TCT Ser	CAG Gln	CCA Pro	aaa Lys 295	CGT Arg	GAC Asp	TTG Leu	GAA Glu	CTG Leu 300	GCA Ala	GTC Val	aag Lys	TIG Leu	10	880
	TTG Leu 305	GAG Glu	AAA Lys	AAG Lys	AGC Ser	CGG Arg 310	AGC Ser	TTT Phe	TTT Phe	GTT Val	GCC Ala 315	TCG Ser	GCT Ala	GGA Gly	TTT Phe	CCT Pro 320	11	L36
15	AGC Ser	GAA Glu	GTT Val	AGG Arg	GAG Glu 325	AGG Arg	CTG Leu	GTT Val	GGA Gly	CTA Leu 330	TAC Tyr	GCA Ala	TTC Phe	TGC Cys	CGG Arg 335	(STC	11	L84
20	ACT Thr	GAT Asp	gat Asp	CIT Leu 340	ATC Ile	GAC Asp	TCT Ser	CCT Pro	GAA Glu 345	GTA Val	TCT Ser	TCC Ser	AAC Asn	CCG Pro 350	CAT His	GCC Ala	12	232
25	ACA Thr	ATT Ile	GAC Asp 355	ATG Met	GTC Val	TCC Ser	gat Asp	TTT Phe 360	CTT Leu	ACC Thr	CTA Leu	CTA Leu	TTT Phe 365	GGG Gly	CCC Pro	CCG Pro	1.2	280
30	Leu	CAC His 370	Pro	Ser	Gln	Pro	<b>Аз</b> р 375	Lys	Ile	Leu	Ser	Ser 380	Pro	Leu	Leu	Pro	13	328
35	385	TCG Ser	His	Pro	Ser	Arg 390	Pro	Thr	Gly	Met	Tyr 395	Pro	Leu	Pro	Pro	Pro 400	13	376
	Pro	TCG Ser	Leu	Ser	Pro 405	Ala	Glu	Leu	Val	Gln 410	Phe	Leu	Thr	Glu	Arg 415	Val	14	124
40	CCC Pro	GTT Val	CAA Gln	TAC Tyr 420	CAT His	TIC Phe	GCC Ala	TTC Phe	AGG Arg 425	TTG Leu	CTC Leu	GCT Ala	AAG Lys	TTG Leu 430	CAA Gln	GGG Gly	14	172
45	CIG Leu	ATC	CCT Pro 435	CGA Arg	TAC Tyr	CCA Pro	CIC	GAC Asp 440	GAA Glu	CTC Leu	CTT Leu	AGA Arg	GGA Gly 445	TAC Tyr	ACC Thir	ACT Thr	15	520
50	gat Asp	CIT Leu 450	ATC Ile	TTT Phe	ccc Pro	TIA Leu	TCG Ser 455	ACA Thr	GAG Glu	GCA Ala	GTC Val	CAG Gln 460	GCT Ala	COG Arg	AAG Lys	ACG Thr	15	568
55	Pro 465	ATC Ile	GAG Glu	ACC Thr	ACA Thr	GCT Ala 470	GAC Asp	TTG Leu	CIG Leu	GAC Asp	TAT Tyr 475	GGT Gly	CTA Leu	TGT Cys	GIA Val	GCA Ala 480	16	516
	GGC Gly	TCA Ser	GIC Val	GCC Ala	GAG Glu 485	CTA Leu	TTG Leu	GTC Val	TAT Tyr	GIC Val 490	TCT Ser	TGG Trp	GCA Ala	AGT Ser	GCA Ala 495	CCA Pro	16	564
60	AGT Ser	CAG Gln	GTC Val	CCT Pro 500	GCC Ala	ACC	ATA Ile	GAA Glu	GAA Glu 505	AGA Arg	GAA Glu	GCT Ala	GIG Val	TTA Leu 510	GTG Val	GCA Ala	1	712
65	AGC Ser	CGA Arg	GAG Glu 515	ATG Met	GGA Gly	ACT	GCC Ala	CTT Leu 520	Gln	TIG Leu	GIG Val	AAC Asn	ATT Ile 525	GCT Ala	AGG Arg	GAC Asp	1	760
<del>7</del> 0	ATT	AAA Lys 530	GGG Gly	gac Asp	GCA Ala	ACA Thr	GAA Glu 535	GG Gly	AGA Arg	TTT Phe	TAC Tyr	CTA Leu 540	Pro	CIC	TCA Ser	TTC Phe	1	808

	TTT Phe 545	GGT Gly	CIT Leu	CGG Arg	gat Asp	GAA Glu 550	TCA Ser	aag Lys	CTT Leu	GCG Ala	ATC Ile 555	CCG Pro	ACT Thr	GAT Asp	TGG Trp	ACG Thr 560	1856
5	GAA Glu	CCT Pro	CGG Arg	CCT Pro	CAA Gln 565	gat Asp	TTC Phe	GAC Asp	AAA Lys	CTC Leu 570	CIC Leu	AGT Ser	CTA Leu	TCT Ser	CCT Pro 575	TCG Ser	1904
10	TCC Ser	ACA Thr	TIA Leu	CCA Pro 580	TCT Ser	TCA Ser	AAC Asn	GCC Ala	TCA Ser 585	GAA Glu	AGC Ser	TTC Phe	CCG Arg	TTC Phe 590	GAA Glu	TGG Trp	1952
15	AAG Lys	ACG Thr	TAC Tyr 595	TCG Ser	CTT Leu	CCA Pro	TTA Leu	GTC Val 600	GCC Ala	TAC Tyr	GCA Ala	GAG Glu	GAT Asp 605	CTT Leu	GCC Ala	AAA Lys	2000
סי	CAT His	TCT Ser 610	TAT Tyr	AAG Lys	GGA Gly	ATT Ile	GAC Asp 615	CGA Arg	CTT Leu	CCT Pro	ACC Thr	GAG Glu 620	GTT Val	CAA Gln	GCG Ala	GGA Gly	2048
<u>.</u>	ATG Met 625	CGA Arg	GCG Ala	GCT Ala	TGC Cys	GCG Ala 630	AGC Ser	TAC Tyr	CTA Leu	CIG Leu	ATC Ile 635	GC Gly	CGA Arg	GAG Glu	ATC Ile	AAA Lys 640	2096
	GTC Val	AT CLL	TGG Trp	AAA Lys	GGA Gly 645	GAC Asp	GTC Val	GGA Gly	GAG Glu	AGA Arg 650	AGG Arg	ACA Thr	GTT Val	GCC Ala	GGA Gly 655	TGG Trp	2144
<b>)</b> 0	ACG Arg	AGA Arg	GTA Val	CGG Arg 660	aaa Lys	GTC Val	TTG Leu	AGT Ser	GIG Val 665	GTC Val	ATG Met	AGC Ser	GGA Gly	TGG Trp 670	GAA Glu	GGG	2192
) \$	GJn CAG	TAAC	ACAC	ecc c	EDAGE	ATAC	C GA	ACAG?	CAAT	CAI	GAGI	GAG	AATZ	raaa/	TCA		2245
	1001	CAAI	cr 1	CITI	cici	DA GO	TGCI	CTT	TT	GITI	TCT	ATT7	ATGAC	CA A	vara:	TAAAGG	2305
<b>K</b> U	AACI	recc	TT	CAGA	TATI	מד כז	CIIC		CAI	CIIC	CIC	CTT	CCAI	rcg 1	TIGI	TCTT	2365
	CCAT	TTT	GT (	XGT1	TACI	TA TO	TCAP	ATTCI	rr	TCIT	GCT	TTT	CIT	ATC F	ATC	AGACA	2425

- (2) INFORMATION FOR SEQ ID NO:19:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 673 amino acids
    - (B) TYPE: amino acid

ATTCIATAGA TGITTAGAAT TTATACAAAA AAAAAAAAA AAAAA

- (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:19:

Met Thr Ala Leu Ala Tyr Tyr Gln Ile His Leu Ile Tyr Thr Leu Pro 1 5 10 15 2470

Ile Leu Gly Leu Leu Thr Ser Pro Ile Leu Thr Lys Phe 20 25 30

Asp Ile Tyr Lys Ile Ser Ile Leu Val Phe Ile Ala Phe Ser Ala Thr 35 40 45

Thr Pro Trp Asp Ser Trp Ile Ile Arg Asn Gly Ala Trp Thr Tyr Pro 50 55 60

Ser Ala Glu Ser Gly Gln Gly Val Phe Gly Thr Phe Leu Asp Val Pro

# SUBSTITUTE SHEET (RULE 26)

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	65					70					75					80
	Tyr	Glu	Glu	Tyr	Ala 85	Phe	Phe	Val	Ile	Gln 90	Thr	Val	Ile	Thr	Gly 95	Leu
5	Val	Tyr	Val	Leu 100	Ala	Thr	Arg	His	Leu 105	Leu	Pro	Ser	Leu	Ala 110	Leu	Pro
10	Lys	Thr	Arg 115	Ser	Ser	Ala	Leu	Ser 120	Leu	Ala	Leu	Lys	Ala 125	Leu	Ile	Pro
	Leu	Pro 130	Ile	Ile	Tyr	Leu	Phe 135	Thr	Ala	His	Pro	Ser 140	Pro	Ser	Pro	Asp
15	Pro 145	Leu	Val	Thr	Asp	His 150	Tyr	Phe	Tyr	Met	Arg 155	Ala	Leu	Ser	Leu	Leu 160
20	Ile	Thr	Pro	Pro	Thr 165	Met	Leu	Leu	Ala	Ala 170		Ser	Gly	Glu	Tyr 175	Ala
20	Phe	Asp	Trp	Lys 180	Ser	Gly	Arg	Ala	Lys 185		Thr	Ile	Ala	Ala 190	Ile	Met
25	Ile	Pro	Thr 195	Val	Tyr	Leu	Ile	Trp 200		Asp	Tyr	Val	Ala 205	Val	Gly	Gln
	Asp	Ser 210	-	Ser	Ile	Asn	Asp 215		Lys	Ile	· Val	Gly 220	_	Arg	Leu	Gly
30	Gly 225	Val	Leu	Pro	Ile	Glu 230		Ala	Met	. Phe	235		Leu	Thr	Asm	Leu 240
35	Met	Ile	· Val	. Leu	Gly 245		Ser	Ala	Cys	Asp 250		Thr	Gln	Ala	Leu 255	Tyr
-	Leu	Leu	His	260		Thr	Ile	тут	Gl _y 265		ı Lys	Lys	: Met	270		Ser
40	Phe	Pro	275		Thr	Pro	Pro	Val 280		ı Se	r Lei	ı Phe	285		Ser	Arg
	Pro	Туз 290		c Ser	Glm	Pro	295		y Asj	Le	u Gli	1 Let 300		a Val	. Lys	Leu
45	Leu 305		ı Ly:	s Lys	s Ser	310		r Phe	e Ph	e Va	1 Ala 31		c Ala	a Gly	Phe	2 Pro 320
50				•	325	5				33	0				33!	
				34	0				34	5				350	)	s Ala
55			35	5				36	0				36	5		o Pro
		37	0				37	5				38	0			u Pro
60	38	5				39	0				39	95				0 Pro 400
65					40	5				4	10				41	
				42	20				4	25				43	30	in Gly
70	Le	u I		ro Ai 35	rg T)	nr Pi	n L		ട്ടോ G 40	lu L	eu L	eu A		ly T) 45	/r Ti	ar Tha

Asp Leu Ile Phe Pro Leu Ser Thr Glu Ala Val Gln Ala Arg Lys Thr 455 Pro Ile Glu Thr Thr Ala Asp Leu Leu Asp Tyr Gly Leu Cys Val Ala 475 Gly Ser Val Ala Glu Leu Leu Val Tyr Val Ser Trp Ala Ser Ala Pro Ser Gln Val Pro Ala Thr Ile Glu Glu Arg Glu Ala Val Leu Val Ala 505 Ser Arg Glu Met Gly Thr Ala Leu Gln Leu Val Asn Ile Ala Arg Asp Ile Lys Gly Asp Ala Thr Glu Gly Arg Phe Tyr Leu Pro Leu Ser Phe Phe Gly Leu Arg Asp Glu Ser Lys Leu Ala Ile Pro Thr Asp Trp Thr 550 Glu Pro Arg Pro Gln Asp Phe Asp Lys Leu Leu Ser Leu Ser Pro Ser Ser Thr Leu Pro Ser Ser Asn Ala Ser Glu Ser Phe Arg Phe Glu Trp 585 Lys Thr Tyr Ser Leu Pro Leu Val Ala Tyr Ala Glu Asp Leu Ala Lys 30 His Ser Tyr Lys Gly Ile Asp Arg Leu Pro Thr Glu Val Gln Ala Gly Met Arg Ala Ala Cys Ala Ser Tyr Leu Leu Ile Gly Arg Glu Ile Lys Val Val Trp Lys Gly Asp Val Gly Glu Arg Arg Thr Val Ala Gly Trp Arg Arg Val Arg Lys Val Leu Ser Val Val Met Ser Gly Trp Glu Gly 665 Gln

(2) INFORMATION FOR SEQ ID NO:20:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 1165 base pairs

(B) TYPE: nucleic acid

(C) STRANDEDNESS: double

(D) TOPOLOGY: linear

55 (ii) MOLECULE TYPE: cDNA

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(iii) HYPOTHETICAL: NO

(iv) ANTI-SENSE: NO

(vi) ORIGINAL SOURCE:

(A) ORGANISM: Phaffia rhodozyma

(ix) FEATURE:

(A) NAME/KEY: CDS

(B) LOCATION: 141..896

(D) OTHER INFORMATION: /product= "PRidi"

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:20:

	CITY	TCT	TTC (	TOG/	CCI	TT	:GGCZ	AGGC(	GT.	rgaac	ACT	cgm	TIAC:	CA :	racco	CACAT	60
	CIC	CATZ	ATA :	ICAC:	TTC	T C	TTC	'AGA	A CA	GIIC	TIGA	GIC	AACC	AA.	AAGAJ	AAGAAG	120
5	GCAC	PAAGI	AA.	PATAI	TCIZ	AG AM Me	NG TO et Se 1	C A	rg Co	C A	AC AT an 11	M GI Le Va	T C	CC CC CP CP:	no Al	CC la 10	170
10	GAG Glu	GTC Val	CGA Arg	ACC Thr	GAA Glu 15	GGA Gly	CTC Leu	AGT Ser	TTA Leu	GAA Glu 20	GAG Glu	TAC Tyr	GAT Asp	GAG Glu	GAG Glu 25	CAG Gln	218
15	Val	AGG Arg	CTG Leu	ATG Met 30	GAG Glu	GAG Glu	CGA Arg	TGT Cys	ATT Ile 35	CIT Leu	GTT Val	AAC Asn	CCG Pro	GAC Asp 40	GAT Asp	GTG Val	266
<b>X</b> 0	GCC Ala	TAT Tyr	GGA Gly 45	GAG Glu	GCT Ala	TCG Ser	aaa Lys	AAG Lys 50	ACC Thr	TGC Cys	CAC His	TIG Leu	ATG Met 55	TCC Ser	AAC Asn	ATC Ile	314
	AAC Astri	OCG Ala 60	CCC Pro	AAG Lys	GAC Asp	CTC Leu	CTC Leu 65	CAC His	CGA Arg	GCA Ala	TTC Phe	TCC Ser 70	GTG Val	TTT Phe	CIC Leu	TTC Phe	362
ש	CCC Arg 75	CCA Pro	TCG Ser	GAC Asp	GGA Gly	GCA Ala 80	CTC Leu	CIG Leu	CTT Leu	CAG Gln	CGA Arg 85	AGA Arg	GCG Ala	GAC Asp	GAG Glu	AAG Lys 90	410
<b>W</b> 2	ATT	ACG Thir	TTC Phe	CCT Pro	GGA Gly 95	ATG Met	TGG Trp	ACC Thr	AAC Asn	ACG Thr 100	TGT Cys	TGC Cys	AGT Ser	CAT His	CCT Pro 105	TIG Leu	458
35	AGC Ser	ATC Ile	AAG Lys	GGC Gly 110	GAG Glu	GIT Val	GAA Glu	GAG Glu	GAG Glu 115	AAC Asn	CAG Gln	ATC Ile	GGT Gly	GTT Val 120	CGA Arg	CGA Arg	506
40	GCT Ala	GCG Ala	TCC Ser 125	CGA Arg	AAG Lys	TIG Leu	GAG Glu	CAC His 130	GAG Glu	CTT Leu	GGC Gly	GIG Val	CCT Pro 135	ACA Thr	TCG Ser	TCG Ser	554
	ACT Thr	CCG Pro 140	CCC Pro	GAC Asp	TCG Ser	TTC Phe	ACC Thr 145	TAC Tyr	CTC Leu	ACT Thr	AGG Arg	ATA Ile 150	CAT His	TAC Tyr	CTC Leu	GCT Ala	602
1,	CCG Pro 155	AGT Ser	GAC Asp	GGA Gly	CTC Leu	TGG Trp 160	GGA Gly	GAA Glu	CAC His	GAG Glu	ATC Ile 165	GAC Asp	TAC Tyr	ATT Ile	CTC Leu	TTC Phe 170	650
50	TCA Ser	ACC Thr	ACA Thr	CCT Pro	ACA Thr 175	GAA Glu	CAC His	ACT Thr	GGA Gly	AAC Asn 180	CCT Pro	AAC Asn	GAA Glu	GTC Val	TCT Ser 185	GAC Asp	698
55	ACT Thr	CGA Arg	TAT Tyr	GTC Val 190	ACC Thr	AAG Lys	ccc Pro	GAG Glu	CIC Leu 195	CAG Gln	GCG Ala	ATG Met	TTT Phe	GAG Glu 200	GAC Asp	GAG Glu	746
60	TCT Ser	AAC Asn	TCA Ser 205	TTT Phe	ACC Thr	CCT	TGG Trp	TTC Phe 210	AAA Lys	TIG Leu	ATT Ile	GCC Ala	CGA Arg 215	GAC Asp	TTC Phe	CTG Leu	794
	TTT Phe	GGC Gly 220	TGG Trp	TGG Trp	GAT Asp	CAA Gln	CIT Leu 225	CTC Leu	GCC Ala	AGA Arg	CGA Arg	AAT Asn 230	GAA Glu	AAG Lys	GGT Gly	GAG Glu	842
65	GTC Val 235	GAT Asp	GCC Ala	AAA Lys	TCG Ser	TIG Leu 240	GAG Glu	GAT Asp	CIC Leu	TCG Ser	GAC Asp 245	AAC Asn	AAA Lys	GTC Val	TGG Trp	AAG Lys 250	890
<b>7</b> 0	ATG	TAG	TCGA		FICT	TICI	GT A	CAGI	CATC	T CA	GITO	GCCT	GIT	GGTT	GCT		943

TGCTTCTTGC TCTTCTTTCT ATATATCTTT TTTCTTGCCT GGGTAGACTT GATCTTTCTA

CATAGCATAC GCATACATAC ATAAACTCTA TTTCTTGTTC TTTATCTCTC TTCTAAGGGA

ATCTTCAAGA TCAATTTCTT TTTGGGCTAC AACATTTCAG ATCAATATTG CTTTTCAGAC

TACAAAAAAA AAAAAAAAA ACTCGAGGGG GGGCCCCGGTA CC

1165

#### (2) INFORMATION FOR SEQ ID NO:21:

15

35

#### (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 251 amino acids
- (B) TYPE: amino acid
- (D) TOPOLOGY: linear

## (ii) MOLECULE TYPE: protein

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO:21:

Met Ser Met Pro Asn Ile Val Pro Pro Ala Glu Val Arg Thr Glu Gly
1 5 10 15

Leu Ser Leu Glu Glu Tyr Asp Glu Glu Glu Glu Val Arg Leu Met Glu Glu 20 25 30

Arg Cys Ile Leu Val Asn Pro Asp Asp Val Ala Tyr Gly Glu Ala Ser 35 40 45

Lys Lys Thr Cys His Leu Met Ser Asn Ile Asn Ala Pro Lys Asp Leu 50 55 60

Leu His Arg Ala Phe Ser Val Phe Leu Phe Arg Pro Ser Asp Gly Ala 65 70 75 80

Leu Leu Gln Arg Arg Ala Asp Glu Lys Ile Thr Phe Pro Gly Met 85 90 95

Trp Thr Asn Thr Cys Cys Ser His Pro Leu Ser Ile Lys Gly Glu Val 100 105 110

Glu Glu Asn Gln Ile Gly Val Arg Arg Ala Ala Ser Arg Lys Leu 115 120 125

Glu His Glu Leu Gly Val Pro Thr Ser Ser Thr Pro Pro Asp Ser Phe 130 135 140

Thr Tyr Leu Thr Arg Ile His Tyr Leu Ala Pro Ser Asp Gly Leu Trp 145 150 155 160

Gly Glu His Glu Ile Asp Tyr Ile Leu Phe Ser Thr Thr Pro Thr Glu 165 170 175

His Thr Gly Asn Pro Asn Glu Val Ser Asp Thr Arg Tyr Val Thr Lys 180 185 190

Pro Glu Leu Gln Ala Met Phe Glu Asp Glu Ser Asn Ser Phe Thr Pro 195 200 205

ω Trp Phe Lys Leu Ile Ala Arg Asp Phe Leu Phe Gly Trp Trp Asp Gln 210 215 220

Leu Leu Ala Arg Arg Asn Glu Lys Gly Glu Val Asp Ala Lys Ser Leu 225 230 235 240

Glu Asp Leu Ser Asp Asn Lys Val Trp Lys Met

	(2) INFO	RMATION FOR SEQ ID NO:22:
5	(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 3550 base pairs  (B) TYPE: nucleic acid  (C) STRANDEINESS: double  (D) TOPOLOGY: linear
10	(ii)	MOLECULE TYPE: DNA (genomic)
	(iii)	HYPOTHETICAL: NO
	(iv)	ANTI-SENSE: NO
15	(vi)	ORIGINAL SOURCE: (A) ORGANISM: Phaffia rhodozyma (B) STRAIN: CBS 6938
20	(ix)	FEATURE: (A) NAME/KEY: exon (B) LOCATION: 941966
25	(xi)	FEATURE: (A) NAME/KEY: intron (B) LOCATION: 9671077
	(xi)	FEATURE: (A) NAME/KEY: excn (B) LOCATION: 10781284
30	(ix)	FEATURE: (A) NAME/KEY: intron (B) LOCATION: 12851364
35	(ix)	FEATURE: (A) NAME/KEY: excen (B) LOCATION: 13651877
40	(ix)	FEATURE: (A) NAME/KEY: intron (B) LOCATION: 18781959
45	(ix	) FEATURE: (A) NAME/KEY: excn (B) LOCATION: 19602202
50	(ix	) FEATURE: (A) NAME/KEY: intron (B) LOCATION: 22032292
<i>3</i> 0	(ix	) FEATURE: (A) NAME/KEY: excm (B) LOCATION: 22933325
55	(ix	) FEATURE: (A) NAME/KEY: CDS (B) LOCATION: join(941966, 10781284, 13651877, 19602202 22933325)
60	(aci	(D) OTHER INFORMATION: /product= "PRGcrtB GB"
		.) SEQUENCE DESCRIPTION: SEQ ID NO:22:  XCAG TITTOCCTTT GACGAGAAG GACACTGGGT TGGAAAGAGA AGATGGTACG 60
હ		COCA CCTTGAATGT GTTGCTTACT AGACATGTTT GACACGCTAA TGCATTTCTT 120
	TOCACI	THEA CITTIGAACT ATGGIGGITG GGGGATCCCC AAAATCATTA GCTTCIACTT 18
<b>7</b> 0	CAN CHARTE	عملت كالمنتاسين علىستشاهات كالمنتاسين كالمنتاسين عميلينين عالمنتاسين على عميل

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								71										
	CCT Pro	TTA Leu	CTT Leu	CCT Pro	CCT Pro 385	TCG Ser	CAC His	CCT Pro	TCC Ser	CGA Arg 390	CCC Pro	ACG Thr	GGA Gly	ATG Met	TAT Tyr 395	CCC Pro	249	91
5	CTC Leu	CCG Pro	CCT Pro	CCT Pro 400	CCT Pro	TCG Ser	CTC Leu	TCG Ser	CCT Pro 405	GCC Ala	GAG Glu	CTC Leu	GTT Val	CAA Gln 410	TTC Phe	CIT Leu	253	39
10	Thr	Glu	Arg 415	Val	Pro	Val	Gln	Tyr 420	CAT His	Phe	Ala	Phe	Arg 425	Leu	Leu	Ala	258	37
15	AAG Lys	TIG Leu 430	CAA Gln	GCG	CIG Leu	ATC	CCT Pro 435	CGA Arg	TAC Tyr	CCA Pro	CTC Leu	GAC Asp 440	GAA Glu	CTC Leu	CTT Leu	AGA Arg	263	15
20	GGA Gly 445	TAC Tyr	ACC Thr	ACT Thr	GAT Asp	CTT Leu 450	ATC Ile	TTT Phe	CCC Pro	TTA Leu	TCG Ser 455	ACA Thr	GAG Glu	GCA Ala	GTC Val	CAG Gln 460	268	13
	GCT Ala	CGG Arg	AAG Lys	ACG Thr	CCT Pro 465	ATC Ile	GAG Glu	ACC Thr	ACA Thr	GCT Ala 470	GAC Asp	TIG Leu	CTG Leu	GAC Asp	TAT Tyr 475	GGT Gly	273	1
ಚ	CIA Leu	TGT Cys	GTA Val	GCA Ala 480	GCC Gly	TCA Ser	GTC Val	GCC Ala	GAG Glu 485	CIA Leu	TTG Leu	GTC Val	TAT Tyr	GTC Val 490	TCT Ser	TGG Trp	277	9
30	GCA Ala	AGT Ser	GCA Ala 495	CCA Pro	AGT Ser	CAG Gln	GTC Val	CCT Pro 500	GCC Ala	ACC Thr	ATA Ile	GAA Glu	GAA Glu 505	AGA Arg	GAA Glu	GCT Ala	282	:7
35	GIG Val	TTA Leu 510	GTG Val	GCA Ala	AGC Ser	CGA Arg	GAG Glu 515	ATG Met	GGA Gly	ACT Thr	GCC Ala	CTT Leu 520	CAG Gln	TIG Leu	GTG Val	AAC Asn	287	5
40	ATT Ile 525	GCT Ala	AGG Arg	GAC Asp	ATT Ile	AAA Lys 530	GGG Gly	GAC Asp	GCA Ala	ACA Thr	GAA Glu 535	GGG Gly	AGA Arg	TTT Phe	TAC Tyr	CTA Leu 540	292	:3
45	CCA Pro	CIC Leu	TCA Ser	TTC Phe	TTT Phe 545	GT Gly	CIT Leu	CGG Arg	GAT Asp	GAA Glu 550	TCA Ser	AAG Lys	CIT Leu	GCG Ala	ATC Ile 555	CCG Pro	297	'1
	ACT Thr	GAT QZA	TGG Trp	ACG Thr 560	GAA Glu	CCT Pro	CGG Arg	CCT Pro	CAA Gln 565	GAT Asp	TTC Phe	GAC Asp	AAA Lys	CIC Leu 570	CIC Leu	AGT Ser	301	.9
50	CTA Leu	TCT Ser	CCT Pro 575	TCG Ser	TCC Ser	ACA Thr	TTA Leu	CCA Pro 580	TCT Ser	TCA Ser	AAC Asn	GCC Ala	TCA Ser 585	GAA Glu	AGC Ser	TTC Phe	306	7
55	CGG Arg	TTC Phe 590	GAA Glu	TGG Trp	AAG Lys	ACG Thr	TAC Tyr 595	TCG Ser	CIT Leu	CCA Pro	TTA Leu	GTC Val 600	GCC Ala	TAC Tyr	GCA Ala	GAG Glu	311	.5
60	GAT Asp 605	CTT Leu	GCC Ala	aaa Lys	CAT His	TCT Ser 610	TAT Tyr	AAG Lys	GGA Gly	ATT Ile	GAC Asp 615	CGA Arg	CTT Leu	CCT Pro	ACC Thr	GAG Glu 620	316	3
65	GTT Val	CAA Gln	GCG Ala	GGA Gly	ATG Met 625	CGA Arg	GCG Ala	GCT Ala	TGC Cys	GCG Ala 630	AGC Ser	TAC Tyr	CTA Leu	CIG Leu	ATC Ile 635	GGC Gly	321	.1
	CGA Arg	GAG Glu	ATC Ile	AAA Lys 640	GTC Val	GIT Val	TGG Trp	AAA Lys	GGA Gly 645	GAC Asp	GIC Val	GGA Gly	GAG Glu	AGA Arg 650	AGG Ar <del>g</del>	ACA Thr	325	9
70	GTT	GCC	GGA	TGG	AGG	AGA	GTA	CGG	AAA	GIC	TIG	AGT	GIG	GIC	ATG	AGC	330	7

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	Val Ala Gly Trp Arg Arg Val Arg Lys Val Leu Ser Val Val Met Ser 655 660 665	
5	GGA TGG GAA GGG CAG TAAGACAGCG GAAGAATACC GACAGACAAT GATGAGTGAG Gly Trp Glu Gly Gln 670	3362
	AATAAAATCA TOOTCAATOT TOTTTOTOTA GGTGCTCTTT TITGTTTTCT ATTATGACCA	3422
0	ACTICIAAAGG AACTGGCCTT GCAGATATTT CTCTTCCCCC CATCTTCCTC CTTTCCATCG	3482
	TITIGITCITT CCATTITIGI CGGITIACIA TGICAATICI TITICITGCI TITICITATC	3542
s	AATCTAGA	3550
	(2) INFORMATION FOR SEQ ID NO:23:	
0	(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 673 amino acids  (B) TYPE: amino acid  (D) TOPOLOGY: linear	
5	(ii) MOLECULE TYPE: protein	
-	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:23:	
ю	Met Thr Ala Leu Ala Tyr Tyr Gln Ile His Leu Ile Tyr Thr Leu Pro 1 5 10 15	
	Ile Leu Gly Leu Leu Gly Leu Leu Thr Ser Pro Ile Leu Thr Lys Phe 20 25 30	
35	Asp Ile Tyr Lys Ile Ser Ile Leu Val Phe Ile Ala Phe Ser Ala Thr 35 40 45	
	Thr Pro Trp Asp Ser Trp Ile Ile Arg Asn Gly Ala Trp Thr Tyr Pro 50 55 60	
40	Ser Ala Glu Ser Gly Gln Gly Val Phe Gly Thr Phe Leu Asp Val Pro 65 70 75 80	
45	Tyr Glu Glu Tyr Ala Phe Phe Val Ile Gln Thr Val Ile Thr Gly Leu 85 90 95	
<b>-</b>	Val Tyr Val Leu Ala Thr Arg His Leu Leu Pro Ser Leu Ala Leu Pro 100 105 110	-
50	Lys Thr Arg Ser Ser Ala Leu Ser Leu Ala Leu Lys Ala Leu Ile Pro 115 120 125	
	Leu Pro Ile Ile Tyr Leu Phe Thr Ala His Pro Ser Pro Ser Pro Asp 130 135 140	
55	Pro Leu Val Thr Asp His Tyr Phe Tyr Met Arg Ala Leu Ser Leu Leu 145 150 155 160	
60	Ile Thr Pro Pro Thr Met Leu Leu Ala Ala Leu Ser Gly Glu Tyr Ala 165 170 175	
<b>ω</b>	Phe Asp Trp Lys Ser Gly Arg Ala Lys Ser Thr Ile Ala Ala Ile Met 180 185 190	
63	Ile Pro Thr Val Tyr Leu Ile Trp Val Asp Tyr Val Ala Val Gly Gln 195 200 205	
	Asp Ser Trp Ser Ile Asn Asp Glu Lys Ile Val Gly Trp Arg Leu Gly 210 215 220	
70	Gly Val Leu Pro Ile Glu Glu Ala Met Phe Phe Leu Leu Thr Asn Leu	

235

230

225

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Met Ile Val Leu Gly Leu Ser Ala Cys Asp His Thr Gln Ala Leu Tyr 245

Leu Leu His Gly Arg Thr Ile Tyr Gly Asn Lys Lys Met Pro Ser Ser

Phe Pro Leu Ile Thr Pro Pro Val Leu Ser Leu Phe Phe Ser Ser Arg

Pro Tyr Ser Ser Gln Pro Lys Arg Asp Leu Glu Leu Ala Val Lys Leu

Leu Glu Lys Lys Ser Arg Ser Phe Phe Val Ala Ser Ala Gly Phe Pro 310

Ser Glu Val Arg Glu Arg Leu Val Gly Tyr Ala Phe Cys Arg Val Thr

Asp Asp Leu Ile Asp Ser Pro Glu Val Ser Ser Asn Pro His Ala Thr

Ile Asp Met Val Ser Asp Phe Leu Thr Leu Leu Phe Gly Pro Pro Leu 360

His Pro Ser Gln Pro Asp Lys Ile Leu Ser Ser Pro Leu Leu Pro Pro

Ser His Pro Ser Arg Pro Thr Gly Met Tyr Pro Leu Pro Pro Pro Pro 390

Ser Leu Ser Pro Ala Glu Leu Val Gln Phe Leu Thr Glu Arg Val Pro 410

Val Gln Tyr His Phe Ala Phe Arg Leu Leu Ala Lys Leu Gln Gly Leu

Ile Pro Arg Tyr Pro Leu Asp Glu Leu Leu Arg Gly Tyr Thr Thr Asp

Leu Ile Phe Pro Leu Ser Thr Glu Ala Val Gln Ala Arg Lys Thr Pro

Ile Glu Thr Thr Ala Asp Leu Leu Asp Tyr Gly Leu Cys Val Ala Gly

Ser Val Ala Glu Leu Leu Val Tyr Val Ser Trp Ala Ser Ala Pro Ser 490

Gln Val Pro Ala Thr Ile Glu Glu Arg Glu Ala Val Leu Val Ala Ser

Arg Glu Met Gly Thr Ala Leu Gln Leu Val Asn Ile Ala Arg Asp Ile

Lys Gly Asp Ala Thr Glu Gly Arg Phe Tyr Leu Pro Leu Ser Phe Phe 535

Gly Leu Arg Asp Glu Ser Lys Leu Ala Ile Pro Thr Asp Trp Thr Glu

Pro Arg Pro Gln Asp Phe Asp Lys Leu Leu Ser Leu Ser Pro Ser Ser

Thr Leu Pro Ser Ser Asn Ala Ser Glu Ser Phe Arg Phe Glu Trp Lys 585

Thr Tyr Ser Leu Pro Leu Val Ala Tyr Ala Glu Asp Leu Ala Lys His

## SUBSTITUTE SHEET (RULE 26)

	Ser	Tyr 610	Ly:	s Gl	y II	e As	p Ar 61	_	u Pr	n Tr	ur G		/al 520	GIn	Ala	G1	у Ми	et			
	Arg 625	Ala	Ala	а Су	rs Al	a Se 63	r Ty 10	r Le	u Le	eu I	_	ly <i>1</i> 35	Arg	Glu	Ile	e Ly		al 40			
	Val	Trp	Ly:	s Gl	y As 64		ıl Gl	.y G)	u Ai		rg T 50	hr V	Val	Ala	Gly	/ Tr 65		rg			
0	Arg	Val	Ar	g Ly 66		ıl Le	eu Se	er Va		al M 55	et S	er (	Зlу	Trp	Glu 670		уG	ln			
5																					
	(2)	INF	ORM	ATIC	ON FC	OR SI	Юπ	OM C	:24 :												
ю		(i	.) S	(A) (B) (C)	LENC TYPI STRI	FIH: E: DI ANDEI	FACT 570 ucle DNES Y: 1	base ic ae S: de	e pa cid cubl	irs											
25		(ii	L) M	OLE	CULE	TYP:	E: C	DINA.											٠		
		(iii	L) F	łYPO	THET	ICAL	: NO														
30		(iv	r) I	WII	-SEN	SE:	NO														
		(v.	i) (				RCE: M: P	haff	ia r	hode	zym	a									
35		(i:	<b>x)</b> 1	(B)	NAM	ATIC	Y: C N: 2 NFOR	45		/pro	oduc	t= '	'PR	-DNA	10"						
40		(x	i) .	SEQU	ENCE	DES	CRII	PTIO	1: SI	D I	D 100	:24	:								
	AA	ACT	TGG	T 17	ÆTTI	rcga.	C GAC	Met	G G G C W											5	0
45	GG Gl _y	y Ly	G A 's T	icc i	ATC I	ACC ( Thr 1	CTT ( Leu ( 15	GAG ( Glu '	STG ( Val (	GAG Glu	TCT Ser	TCT Ser 20	As	C AC	X A	rc o le A	ASP ASP	AAC Asn 25	1	<u>5</u>	8
50	GT Va	CA4	AG C	la :	AAG 1 Lys :	ATC ( Ile ( 30	CAG ( Gln .	GAC Asp	AAG Lys	GAA Glu	GGA Gly 35	ATI	Pr	C CC	T G	AT ( sp (	CAG Sln 40	CAC Glr	3 1	14	16
55	CG Ar	A C. g L	rr / eu :	ATC Ile	TTC · Phe · 45	GCC Ala	GT Gly	AAG Lys	CAG Gln	CTC Leu 50	GAG Glu	GAT Ast	G G	C O	GA A	CC ( hr : 55	CTT Leu	Sei	g r	1	94
60	GA As	тт ърт	AC I	AAC Asn 60	ATC Ile	CAG Gln	aaa Lys	GAG Glu	TCC Ser 65	ACC Thr	CIC Leu	CA(	C C	IC G eu V	TC C al I 70	TT .eu	AGG Arg	Tr	G u	2	42
65	CX Aı	SAG OGG	GA 1y 75	GGA Gly	GCC Ala	aag Lys	AAG Lys	CGA Arg 80	AAG Lys	AAG Lys	AAG Lys	GL:	n T	AC A yr 1 85	CT I	ACC Ihr	CCC Pro	Ly	G 's	2	90
υ,	Ŀ	AG A ys I 90	TC le	aac Lys	CAC His	AAG Lys	CGA Arg 95	AAG Lys	AAG Lys	GTC Val	AAG Lys	AT Me 10	t A	CT I	Ile	CIT Leu	AAG Lys	TA Ty	x	3	328
20	т	ז אמ	מע	حلات	GAC	بلمك	GAT	GGA	AAG	ATC	: AAC	3 03	A C	TT (	CT	CGA.	GAC	3 10	3C	:	386

											•								
	Тут	Ly:	s Va	l Asp	Ser 110	Asp	Gly	/ Lys	5 Il∈	Ly:	s Arg	J Lei	ı Arç	g Arg	Gl: 120		S		
5	CCC Pro	CA(	G CCC	C CAC O Glr 125	ı cys	Gly	A GCI 7 Ala	GCI Gly	TATO Ile	Phe	ATC Met	GC. Ala	r Tro	CA( His	Sei	AA C Asi	C n		434
10	CGA Arg	A CAC	AC. Thi	r TGC c Cys o	GGA Gly	AAG Lys	TGI Cys	GGI Gly 145	Leu	ACC Thi	Tyr	C ACC	TTC Phe 150	: Ala	GA(	GG/	A Y		482
.,	ACC Thr	CAC Glr 155	PIC	C TCI Ser	GCT Ala	TAG	ATCA	ICA	ATCG	TTT	TT C	CCGA	GOG	ar Ci	TTG	GICI	r		537
15	TIG	TTAC	TTA	CICA	AAAA	AA A	AAAA	AAAA	a aa	A								;	570
30	(2)	INF		TION															
			(1)	(B	) LE ) TY ) TO	NGTH PE:	: 15 amin	8 am	ino a id	: acid	s								
. <del></del>		(	ii)	MOLE	CULE	TYP	E: p:	rote	in										
				SEQU															
10	Met 1	Gln	Ile	Phe	Val 5	Lys	Thr	Leu	Thr	Gly 10	Lys	Thr	Ile	Thr	Leu 15				
35	Val	Glu	Ser	Ser 20	Asp	Thr	Ile	Asp	Asn 25	Val	Lys	Ala	Lys	Ile 30	Gln	Asp			
	Lys	Glu	Gly 35	Ile	Pro	Pro	Asp	Gln 40	Gln	Arg	Leu	Ile	Phe 45	Ala	Gly	Lys			
40	Gln	Leu 50	Glu	Asp	Gly	Arg	Thr 55	Leu	Ser	Asp	Tyr	Asn 60	Ile	Gln	Lys	Glu			
	Ser 65	Thr	Leu	His	Leu	Val 70	Leu	Arg	Leu	Arg	Gly 75	Gly	Ala	Lys	Lys	Arg 80			
45	Lys	Lys	Lys	Gln	Tyr 85	Thr	Thr	Pro	Lys	Lys 90	Ile	Lys	His	Lys	Arg 95	Lys			
50	Lys	Val	Lys	Met 100	Ala	Ile	Leu	Lys	Tyr 105	Tyr	Lys	Val	Asp	Ser 110	Asp	Gly			
	Lys	Ile	Lys 115	Arg	Leu	Arg	Arg	Glu 120	Cys	Pro	Gln	Pro	Gln 125	Cys	Gly	Ala			
<b>5</b> 5	Gly	Ile 130	Phe	Met	Ala	Phe	His 135	Ser	Asn	Arg	Gln	Thr 140	Cys	Gly	Lys	Cys			
	Gly 145	Leu	Thr	Tyr	Thr	Phe 150	Ala	Glu	Gly	Thr	Gln 155	Pro	Ser	Ala					
60	(2)	INFO	ORMA:	MOLI	FOR	SEQ	ID N	<b>1</b> 0:26	i:										
65		(i)	(2 (E (C	QUENC A) LE B) TY C) ST O) TO	NGIH PE: RAND	: 30 nucl EDNE	3 ba eic SS:	se p acid doub	airs I										

# SUBSTITUTE SHEET (RULE 26)

(ii) MOLECULE TYPE: CDNA

55

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	(iii) HYPOTHETICAL: NO	
	(iv) ANTI-SENSE: NO	
s	(vi) ORIGINAL SOURCE: (A) ORGANISM: Phaffia rhodozyma	
10	(ix) FEATURE:  (A) NAME/KEY: CDS  (B) LOCATION: 57278  (D) OTHER INFORMATION: /product= "PRoDNALL"	
15	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:26:	
	TITIACACACA AACCTITACCT ACCTTTTCAA CAACAAATCA CACCTAAGCT TACATC	56
20	ATG GAG TOC ATC AAG ACC TOG ATT TOC AAC GCC GCC AAC TAC GCT TCT Met Glu Ser Ile Lys Thr Ser Ile Ser Asn Ala Ala Asn Tyr Ala Ser 1 5 10 15	104
	GAG ACT GTC AAC CAG GCC ACT AGC GCC ACC TCC AAG GAG GCC AAC AAG Glu Thr Val Asm Gln Ala Thr Ser Ala Thr Ser Lys Glu Ala Asm Lys 20 25 30	152
25	GAG GTT GCC AAG GAC TCC AAT GCC GGA GTT GGA ACC CGA ATC AAC GCC Glu Val Ala Lys Asp Ser Asn Ala Gly Val Gly Thr Arg Ile Asn Ala 35 40 45	200
30		
	GCA ATT GAT GCT CTT GGA GAC AAG GCC GAC GAG ACT TCG TCT GAT GCC Gly Ile Asp Ala Leu Gly Asp Lys Ala Asp Glu Thr Ser Ser Asp Ala 50 55 60	248
35	AAG TOC AAG GOC TAC AAG CAG AAC ATC TAAGITATIT AGATAGIOGI Lys Ser Lys Ala Tyr Lys Gln Asn Ile 65 70	295
40	CCATATTT	303
45	(2) INFORMATION FOR SEQ ID NO:27:  (i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 73 amino acids  (B) TYPE: amino acid  (D) TOPOLOGY: linear	
50	(ii) MOLECULE TYPE: protein	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:27:	
55	Met Glu Ser Ile Lys Thr Ser Ile Ser Asn Ala Ala Asn Tyr Ala Ser 1 5 10 15	
	Glu Thr Val Asm Gln Ala Thr Ser Ala Thr Ser Lys Glu Ala Asm Lys 20 25 30	
60	Glu Val Ala Lys Asp Ser Asn Ala Gly Val Gly Thr Arg Ile Asn Ala 35 40 45	
65	Gly Ile Asp Ala Leu Gly Asp Lys Ala Asp Glu Thr Ser Ser Asp Ala 50 55 60	
	Lys Ser Lys Ala Tyr Lys Gln Asn Ile 65 70	

70 (2) INFORMATION FOR SEQ ID NO:28:

5	(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 307 base pairs  (B) TYPE: nucleic acid  (C) STRANDEINESS: double  (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: CDNA	
10	(iii) HYPOTHETICAL: NO	
10	(iv) ANTI-SENSE: NO	
15	(vi) ORIGINAL SOURCE: (A) ORGANISM: Phaffia rhodozyma	
	(ix) FEATURE: (A) NAME/KEY: CDS	
20	(B) LOCATION: 3227 (D) OTHER INFORMATION: /product= "PRCINAL8"	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:28:	
25	AC CCT TCC ATC GAG TCT GAG GCC CGA CAA CAC AAG CTC AAG AGG CTT Pro Ser Ile Glu Ser Glu Ala Arg Gln His Lys Leu Lys Arg Leu 1 5 10 15	47
30	GTG CAG AGC CCC AAC TCT TTC TTC ATG GAC GTC AAG TGC CCT GGT TGC Val Gln Ser Pro Asn Ser Phe Phe Met Asp Val Lys Cys Pro Gly Cys 20 25 30	95
35	TTC CAG ATC ACC GTG TTC TCG CAC GCT TCC ACT GCC GTT CAG TGT Phe Gln Ile Thr Thr Val Phe Ser His Ala Ser Thr Ala Val Gln Cys 35 40 45	143
40	GGA TCG TGC CAG ACC ATC CTC TGC CAG CCC CGG GGA GGA AAG GCT CGA Gly Ser Cys Gln Thr Ile Leu Cys Gln Pro Arg Gly Gly Lys Ala Arg 50 55 60	191
45	CTT ACC GAG GGA TGC TCT TTC CGA CGA AAG AAC TAAGITTCTG TTATCGGATG Leu Thr Glu Gly Cys Ser Phe Arg Arg Lys Asn 65 70 75	244
	ATGCATTCAA ATAAAAGTCA AAAAAAAAA AAAAAAAAC TCGAGGGGGG GCCCGGTACC	304
	CAA	307
50	(2) INFORMATION FOR SEQ ID NO:29:	
55	(i) SEQUENCE CHARACTERISTICS:  (A) LENGIH: 74 amino acids  (B) TYPE: amino acid  (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: protein	
60	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:29:	
	Pro Ser Ile Glu Ser Glu Ala Arg Gln His Lys Leu Lys Arg Leu Val 1 10 15	
65	Gln Ser Pro Asm Ser Phe Phe Met Asp Val Lys Cys Pro Gly Cys Phe 20 25 30	
70	Gln Ile Thr Thr Val Phe Ser His Ala Ser Thr Ala Val Gln Cys Gly 35 40 45	

	Ser Cys Gln Thr Ile Leu Cys Gln Pro Arg Gly Gly Lys Ala Arg Leu 50 55 60	
,	Thr Glu Gly Cys Ser Phe Arg Arg Lys Asn 65 70	
	(2) INFORMATION FOR SEQ ID NO:30:	
,	(i) SEQUENCE CHARACIERISTICS:  (A) LENGTH: 502 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: double	
5	(D) TOPOLOGY: linear (ii) MOLECULE TYPE: CDNA	
	(iii) HYPOTHETICAL: NO	
D	(iv) ANTI-SENSE: NO	
	(vi) ORIGINAL SOURCE: (A) ORGANISM: Phaffia rhodozyma	
•	(1)X) FEATURE: (A) NAME/KEY: CDS (B) LOCATION: 30359 (D) OTHER INFORMATION: /product= "PRODNA35"	
0	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:30:	
15	GTCASCTCCG GCTTAAATCG ATTCGTACA ATG TCT GAA CTC GCC GCC TCC TAC Met Ser Glu Leu Ala Ala Ser Tyr 1 5	53
	GCC GCT CTT ATC CTC GCC GAC GAG GGT ATT GAG ATC ACC TCT GAG AAG Ala Ala Leu Ile Leu Ala Asp Glu Gly Ile Glu Ile Thr Ser Glu Lys 10 15 20	101
<b>10</b>	CTC GTC ACT CTC ACC ACC GCC GCC AAG GTT GAG CTT GAG CCC ATC TGG Leu Val Thr Leu Thr Thr Ala Ala Lys Val Glu Leu Glu Pro Ile Trp 25 30 35 40	149
45	GCC ACT CTC CTT GCC AAG GCC CTC GAG GGA AAG AAC GTC AAG GAG TTG Ala Thr Leu Leu Ala Lys Ala Leu Glu Gly Lys Asn Val Lys Glu Leu 45 50 55	197
50	CTT TCC AAC GTC GGA TCC GGA GCC GGA GGA GCT GCC CCC GCC GCC GCC Leu Ser Asn Val Gly Ser Gly Ala Gly Gly Ala Ala Pro Ala Ala Ala 60 65 70	245
55	GTC GCC GGT GGA GCT TCC GCT GAC GCC TCT GCC CCC GCT GAG GAG AAG Val Ala Gly Gly Ala Ser Ala Asp Ala Ser Ala Pro Ala Glu Glu Lys 75 80 85	293
	AAG GAG GAG AAG GCT GAG GAC AAG GAG GAG TCT GAC GAC GAC ATG GGT Lys Glu Glu Lys Ala Glu Asp Lys Glu Glu Ser Asp Asp Asp Met Gly 90 95 100	341
60	TTC GGA CIT TTC GAT TAAACTCCCT CGCCTAAAAA CCCTTTTCTT CAACCCCCTC Phe Gly Leu Phe Asp 105 110	396
65	TOGIOGCATO GITCACIOGA COGCIGOGIT TGIIGIOCTI TOCICAOGAA TTIIGIOCTI	456
	GICTOGITIC CCAATNOGAT NICCITGAAA TGANGITICC CAATIG	50:
70	(כ) אוניסטטאאינידוראו וויסט פוויס דדי אוני-פו	

	(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 109 amino acids  (B) TYPE: amino acid  (D) TOPOLOGY: linear	
5	(ii) MOLECULE TYPE: protein	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:31:	
10	Met Ser Glu Leu Ala Ala Ser Tyr Ala Ala Leu Ile Leu Ala Asp Glu 1 5 10 15	
15	Gly Ile Glu Ile Thr Ser Glu Lys Leu Val Thr Leu Thr Thr Ala Ala 20 25 30	
	Lys Val Glu Leu Glu Pro Ile Trp Ala Thr Leu Leu Ala Lys Ala Leu 35 40 45	
20	Glu Gly Lys Asn Val Lys Glu Leu Leu Ser Asn Val Gly Ser Gly Ala 50 55 60	
	Gly Gly Ala Ala Pro Ala Ala Ala Val Ala Gly Gly Ala Ser Ala Asp 65 70 75 80	
25	Ala Ser Ala Pro Ala Glu Glu Lys Lys Glu Glu Lys Ala Glu Asp Lys 85 90 95	
30	Glu Glu Ser Asp Asp Met Gly Phe Gly Leu Phe Asp 100 105	
	(2) INFORMATION FOR SEQ ID NO:32:	
35	(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 381 base pairs  (B) TYPE: nucleic acid  (C) STRANDEINESS: double  (D) TOPOLOGY: linear	
<b>4</b> O	(ii) MOLECULE TYPE: cDNA	
	(iii) HYPOTHETICAL: NO	
45	(iv) ANTI-SENSE: NO	
	(vi) ORIGINAL SOURCE: (A) ORGANISM: Phaffia rhodozyma	
50	(ix) FEATURE:  (A) NAME/KEY: CDS  (B) LOCATION: 7282  (D) OTHER INFORMATION: /product= "PRCDNA38"	
35	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:32:	
	CTCAAG ATG ACC AAA GGT ACC TCC TCT TTC GGT AAG CGA CAC ACC AAG Met Thr Lys Gly Thr Ser Ser Phe Gly Lys Arg His Thr Lys  1 5 10	48
50 -	ACC CAC ACC ATC TGC CGA CGA TGT GGT AAC AGG GCT TTC CAC AGG CAG Thr His Thr Ile Cys Arg Arg Cys Gly Asn Arg Ala Phe His Arg Gln 15 20 25 30	9(
55	AAG AAG ACC TGT GCC CAG TGT GGA TAC CCT GCC GCC AAG ATG CGA AGC Lys Lys Thr Cys Ala Gln Cys Gly Tyr Pro Ala Ala Lys Met Arg Ser 35 40 45	144
<b>7</b> 0	TTC AAC TGG GGA GAG AAG GCC AAG AGG AGA AAG ACC ACC	192

	50	55	60	
5	CGA ATG CAG CAC CTC AAG GAC GT Arg Met Gln His Leu Lys Asp Va 65			240
	CGA GAG GGA ACT TCC GCC ACC AA Arg Glu Gly Thr Ser Ala Thr Ly 80 85			289
0	ATCCATCACC TOGTGATCAG GGCGGGIA	AT AATCTTTGT	TAGAGACTAT CCATGITCIO	G 349
	CTGCCGCATC AAACAAAAAA AAAAAAAA	AA AA		381
5	(2) INFORMATION FOR SEQ ID NO:	:33:		
20	(i) SEQUENCE CHARACTERI (A) LENGTH: 91 an (B) TYPE: amino a (D) TOPOLOGY: lir	mino acids acid		
	(ii) MOLECULE TYPE: prot	tein		
25	(xi) SEQUENCE DESCRIPTION	ON: SEQ ID NO:	:33:	
	Met Thr Lys Gly Thr Ser Ser Pl	ne Gly Lys Arq 10	His Thr Lys Thr His 15	
30	Thr Ile Cys Arg Arg Cys Gly A	sn Arg Ala Phe 25	e His Arg Gln Lys Lys 30	
	Thr Cys Ala Gln Cys Gly Tyr P 35	ro Ala Ala Ly: 40	s Met Arg Ser Phe Asn 45	
35	Trp Gly Glu Lys Ala Lys Arg A 50 55	rg Lys Thr Th	r Gly Thr Gly Arg Met 60	
40	Gln His Leu Lys Asp Val Ser A 65 70	arg Arg Phe Ly 7		
	Gly Thr Ser Ala Thr Lys Lys V 85	al Lys Ala Gl 90	u ·	
45	(2) INFORMATION FOR SEQ ID NO	0:34:		
50	(i) SEQUENCE CHARACTERIS  (A) LENGIH: 473 bas  (B) TYPE: nucleic a  (C) STRANDEDNESS: 0  (D) TOPOLOGY: lines	se pairs acid double		
55	(ii) MOLECULE TYPE: CDNA			
	(iii) HYPOTHETICAL: NO			
60	(iv) ANTI-SENSE: NO (vi) ORIGINAL SOURCE:	<b>.</b>		
65	(A) ORGANISM: Phaf  (ix) FEATURE:  (A) NAME/KEY: CDS  (B) LOCATION: 19  (D) OTHER INFORMAT	321	= "PRCDNA46"	
			24	

(2) INFORMATION FOR SEQ ID NO:36:

5	(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 608 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: double  (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: cDNA	
	(iii) HYPOTHETICAL: NO	
10	(iv) ANTI-SENSE: NO	
	(vi) ORIGINAL SOURCE: (A) ORGANISM: Phaffia rhodozyma	
20	(ix) FEATURE: (A) NAME/KEY: CDS (B) LOCATION: 18453 (D) OTHER INFORMATION: /product= "FRCDNA64"	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:36:	
25	AAGACTCGTC GTTCAGC ATG TCC TCC GTC AAA GCC ACC AAA GGA AAG GGT Met Ser Ser Val Lys Ala Thr Lys Gly Lys Gly 1 5 10	50
30	CCC GCC GCC TCG GCT GAT GTT AAG GCC AAG GCC GCC AAG AAG GCT GCC Pro Ala Ala Ser Ala Asp Val Lys Ala Lys Ala Ala Lys Lys Ala Ala 15 20 25	98
	CTC AAG GGT ACT CAG TCT ACT TCC ACC AGG AAG GTC CGA ACT TCG GTC Leu Lys Gly Thr Gln Ser Thr Ser Thr Arg Lys Val Arg Thr Ser Val 30 35 40	146
35	TCT TTC CAC CGA CCC AAG ACT CTC CGA CTT CCC CGA GCT CCC AAG TAC Ser Phe His Arg Pro Lys Thr Leu Arg Leu Pro Arg Ala Pro Lys Tyr 45 50 55	194
40	CCC CGA AAG TCG GTC CCT CAC GCC CCT CGA ATG GAT GAG TTC CGA ACT Pro Arg Lys Ser Val Pro His Ala Pro Arg Met Asp Glu Phe Arg Thr 60 65 70 75	242
45	ATC ATC CAC CCC TTG GCT ACC GAG TCC GCC ATG AAG AAG ATT GAG GAG Ile Ile His Pro Leu Ala Thr Glu Ser Ala Met Lys Lys Ile Glu Glu 80 85 90	290
50	CAC AAC ACC CTT GTG TTC ATC GTC GAT GTC AAG TCC AAC AAG CGA CAG His Asn Thr Leu Val Phe Ile Val Asp Val Lys Ser Asn Lys Arg Gln 95 100 105	338
	ATC AAG GAC GCC GTC AAG AAG CTC TAC GAG GTC GAT ACC GTC CAC NTC Ile Lys Asp Ala Val Lys Lys Leu Tyr Glu Val Asp Thr Val His Xaa 110 115 120	386
55	AAC NOC TIG ATC ACC COC GOC GGA AGG AAG AAG CTT ACG TOC GAC TTA Asn Xaa Leu Ile Thr Pro Ala Gly Arg Lys Lys Leu Thr Ser Asp Leu 125 130 135	434
60	CCC CCG ACC ACG ACG CTC T TAACGITGCC AACAAGGCCG GCTACATCTA Pro Pro Thr Thr Leu 140 145	483
	ATCEACTCCA TCCCTIGGAT CGGTTCAGTT GTTTGGTTTG CATCCGGTTT CAGAGTTTGA	543
65	CEACCTTGAA ACTONAANAC TTTGGATGCA TGTTTGAAAT TCTCNAAATA AAAAAAAAAA	603
	AAAAA	608

(2) INFORMATION FOR SEQ ID NO:37:

5	(D) TOPOLOGY: linear													
	(ii) MOLECULE TYPE: protein													
10	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:37:													
	Met Ser Ser Val Lys Ala Thr Lys Gly Lys Gly Pro Ala Ala Ser Ala 1 5 10 15													
15	Asp Val Lys Ala Lys Ala Lys Lys Ala Ala Leu Lys Gly Thr Gln 20 25 30													
20	Ser Thr Ser Thr Arg Lys Val Arg Thr Ser Val Ser Phe His Arg Pro 35 40 45													
	Lys Thr Leu Arg Leu Pro Arg Ala Pro Lys Tyr Pro Arg Lys Ser Val 50 55 60													
25	Pro His Ala Pro Arg Met Asp Glu Phe Arg Thr Ile Ile His Pro Leu 65 70 75 80													
	Ala Thr Glu Ser Ala Met Lys Lys Ile Glu Glu His Asn Thr Leu Val 85 90 95													
30	Phe Ile Val Asp Val Lys Ser Asn Lys Arg Gln Ile Lys Asp Ala Val 100 105 110													
35	Lys Lys Leu Tyr Glu Val Asp Thr Val His Xaa Asn Xaa Leu Ile Thr 115 120 125													
	Pro Ala Gly Arg Lys Lys Leu Thr Ser Asp Leu Pro Pro Thr Thr 130 135 140													
40	Leu 145													
	(2) INFORMATION FOR SEQ ID NO:38:													
45	(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 466 base pairs  (B) TYPE: nucleic acid  (C) STRANDEINESS: double  (D) TOPOLOGY: linear													
50	(ii) MOLECULE TYPE: CDNA													
	(iii) HYPOTHETICAL: NO													
55	(iv) ANTI-SENSE: NO													
	(vi) ORIGINAL SOURCE: (A) ORGANISM: Phaffia rhodozyma													
60	(ix) FEATURE:  (A) NAME/KEY: CDS  (B) LOCATION: 81416  (D) OTHER INFORMATION: /product= "PRoDNA68"													
65	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:38:													
	CTTTGAACCT CCAACCTCGG CATCAAGCAC TAGTCAGCCT CGGCTTAAAT CGATTCGTGT	60												
70	AGCCTITCAA ACTCGTAAAA ATG AAG CAC ATC GCC GCT TAC TIG CTC CTC  Met Lys His Ile Ala Ala Tyr Leu Leu Leu	110												

							1				5				:	LO		
5			GGT Gly															158
			ACC Thr															206
10	ATC Ile	TCC Ser	GAG Glu 45	CIT Leu	AAC Asn	GGC Gly	AAG Lys	GAC Asp 50	TTG Leu	AAC Asn	ACC Thr	CTC Leu	ATC Ile 55	Ala	GAG Glu	GGA Gly		254
15			Lys													GCT Ala		302
20							Gly					Ala				AAG Lys 90		350
25	AAG Lys	GAG Glu	GAG Glu	AAG Lys	GIC Val 95	Glu	GAC Asp	AAG Lys	GAC Glu	GAG Glu 100	Ser	GAC Asp	GAC Asp	GAC Asp	ATG Met	Gly		398
			CII Leu		Asp		ACTO	CIT	ACAC	CTTI	TT C	AAAC	TCTI	ന്ദ് ദ	TGG	TOGA		453
30	GGG	GGGG	cac	ŒT														466
35	(2)	IN	ORM (i)	SEQU	ENCE A) LE B) TY	E CHA	RACT H: 11	TERIS	STIC nino cid	_	ìs							
40			(ii)		ECULI													
			(xi)	SEQ	UENC	E DE	SCRI!	PTIO	ท: S	EQ I	D 1/10	:39:						
45		L Ly	s Hi	s Il		a Al	а Ту	r Le	u Le		u Al O	a Th	r Gl	y Gl	y As	n Xaa 5		
	Se	r Pr	o Se		a Al O	a As	p Va	l Ly		a Le 5	u Le	u Al	a Tr		l As	p Ile	:	
50	Gl	u Al	_	p As	p Al	a Ar	g Le		u Tr 0	r Le	eu Il	.e Se		u Le 15	eu As	m Gly	,	
55	-	5	50				5	55				€	50			La Ser		
	6	55				•	70				•	75				la Gly 80	0	
60					8	35				!	90					al Gli 95	u	
65		sp L	ys G		lu Se 00	er A	sp A	sp A		et G .05	TA 5	he G	ly L		he A .10	sp		
	(:	2) I	NFOR	MATI	ON F	OR S	EQ I	D 100	:40:									

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 570 base pairs

			(	(B) I (C) S (D) I	TRAN	DEDIN	ESS:	dou									
5		(ii	) MC	LECU	LE T	YPE:	CDN	ZA.									
		(iii	) HY	POIH	ETIC	AL:	<b>N</b> O										
10		(iv	AA (	m-s	ENSE	: NO	1										
10		(vi		IGIN A) O				ffia	rho	dozy	ma						
15		(ix		ATUR A) N		vev.	CTCC										
			(	B) L D) O	CAT	ION:	49.	.501	: /p	rodu	ct=	"PRC	DINIA7	3"			
20				QUEN													
25	CIT	CCTC	CCG	TCAA	GGCA	AA C	CITC	AGAA'	roc	TCTC	AAGT	CAT	ICAA	Me		A CGA y Arg	57
_	GTC Val	CGC	ACC Thr	AAA Lys	ACC Thr	GIC	AAG	CGA	GCT	TCG	CGA	GIG	ATG	ATC	GAG	AAG	105
		5		2,5		Val	10	Arg	ма	SeI	Arg	15	Met	ше	Glu	Lys	
30	TTC Phe	TAC	CCT	CGA Arg	CIC	ACT	CTT	GAT	TTC	CAC	ACC	AAC	AAG	CGA	ATC	GCC	153
	20	•		5		25		p	1110	1113	30	ASII	ьуs	Arg	TTE	35	
35	GAC GAC	GAG Glu	GTT Val	GCC Ala	ATC Ile 40	ATC Ile	CCC Pro	TCC Ser	AAG Lys	OGA Arg 45	CTT Leu	CGA Arg	AAC Asn	AAG Lys	ATC Ile 50	GCT Ala	201
40	GGG Gly	TTC Phe	ACT Thr	ACC Thr 55	CAC His	TIG Leu	ATG Met	AAG Lys	CGA Arg 60	ATC Ile	CAG Gln	AAG Lys	GGA Gly	CCC Pro 65	GTT Val	CGA Arg	249
	GGT	ATC	TCC	TTC	AAG	CTT	CAG	GAG	GAG	GAG	CGA	GAG	AGG	AAG	GAT	CAG	297
45	GIY	116	70	Phe	Lys	Leu	Gin	75	GIu	Glu	Arg	Glu	Arg 80	Lys	Asp	Gln	
•	TAC Tyr	GTT Val 85	CCT Pro	GAG Glu	GTC Val	TCC Ser	GCC Ala 90	Leu	GCC Ala	GCC Ala	CCT Pro	GAG Glu 95	Leu	GGT Gly	TIG Leu	GAG Glu	345
50	GIT	GAC	œc	GAC	ACC	AAG	GAT	CTT	crc	CGA	TCC	CIT	GGC	ATG	GAC	TCC	393
	Val 100	Asp	Pro	Asp	Thr	Lys 105	Asp	Leu	Leu	Arg	Ser 110	Leu	Gly	Met	Asp	Ser 115	223
55	ATC Ile	AAC Asn	GTC Val	CAG Glin	GTC Val 120	TCC Ser	GCT Ala	CCT Pro	ATC Ile	TCT Ser 125	TCC Ser	TAC Tyr	GCT Ala	GCC Ala	CCC Pro 130	GAG Glu	441
60	CGA Arg	GT Gly	CCC Pro	CGA Arg 135	GT Gly	GCC Ala	GGA Gly	CGA Arg	NGT Xaa 140	GGA Gly	CGA Arg	ATC Ile	GTC Val	CCC Pro 145	GGA Gly	GCT Ala	489
4	GGC Gly	CGA Arg	TAC Tyr 150	TAAC	FIGIT	TT (	770	AACC2	∌n G	<b>SGAT</b> Z	TTT	TA E	TATTY	CCT			538
65	AGG	TTG	AAA :	rryy	TTAT	C A	TCT	LCC13	AT A								570
	(2)	INFC	ORIMA:	rion	FOR	SEQ	ID 1	<b>1</b> 0:43	L:								

			(i) S	(A) (B)	INCE TYP TOP	GIH: Œ: a	150 mino	ami aci	no a		i							
5		(:	ii) N	/OLE	ULE	TYPE	E: pr	otei	n									
		(၁	ki) 9	EEQUE	NCE	DESC	RIPI	MOIT:	SEÇ	D ID	NO:4	11:						
10	Met 1	Gly	Arg	Val	Arg 5	Thr	Lys	Thr	Val	Lys 10	Arg	Ala	Ser	Arg	Val 15	Met		
15	Ile	Glu	Lys	Phe 20	Tyr	Pro	Arg	Leu	Thr 25	Leu	Asp	Phe	His	Thr 30	Asn	Lys		
	Arg	Ile	Ala 35	Asp	Glu	Val	Ala	Ile 40	Ile	Pro	Ser	Lys	Arg 45	Leu	Arg	Asm		
20	Lys	Ile 50		Gly	Phe	Thr	Thr 55	His	Leu	Met	Lys	Arg 60	Ile	Gln	Lys	Gly		
	Pro 65	Val	Arg	Gly	Ile	Ser 70	Phe	Lys	Leu	Gln	Glu 75	Glu	Glu	Arg	Glu	Arg 80		
25	Lys	Asp	Gln	Tyr	Val 85	Pro	Glu	Val	Ser	Ala 90	Leu	Ala	Ala	Pro	Glu 95	Leu		
•	Gly	Leu	Glu	Val 100	Asp	Pro	Asp	Thr	Lys 105	Asp	Leu	Leu	Arg	Ser 110	Leu	Gly		
30	Met	Asp	Ser 115		Asn	Val	Gln	Val 120	Ser	Ala	Pro	Ile	Ser 125		Tyr	Ala		
35	Ala	Pro 130		Arg	Gly	Pro	Arg 135	_	Ala	Gly	Arg	Хаа 140	Gly	Arg	Ile	Val		
	Pro 145	_	Ala	Gly	Arg	Tyr 150												
40	(2)	INF	ORMA	TION	FOR	SEQ	) ID	NO:4	2:									
45		i)	(	(A) I (B) T (C) S	CE C ENGT YPE: TRAN OPOL	H: 3 muc DEDN	73 b leic ESS:	ase aci dou	pair d	s								
50					LE T			IA.										
					ENSE													
55			i) O	RIGI	VAL S	SOURC	Œ:	affia	a rho	odoz	/ma							
		( <b>i</b> :	x) F	EATU														
60				(B)	OTHE	LICM	: 13	324		prod	uct=	"PR	cDNA	76"				
65		(x	i) S	EQUE	NCE 1	DESC	RIPT	ION:	SEQ	110	NO:4	2:						
	cc	ATCA	TCCA		ATG ( Met )													48
70	AA	G AC	C CA	G AA	g aa	g aa	g aa	G TG	G TO	C AA	G GG	A AA	G GI	G AA	G GA	CAA	G	96

	Lys	Thr	Gln 15	Lys	Lys	Lys	Lys	Trp 20	Ser	Lys	Gly	Lys	Val 25	Lys	Asp	Lys		
5	GCC Ala	GCC Ala 30	CAC His	CAC His	GTC Val	GTT Val	GTT Val 35	GAT Asp	CAG Gln	GCC Ala	ACT Thr	TAC Tyr 40	GAC Asp	AAG Lys	ATC Ile	GIT Val		144
10	AAG Lys 45	GAG Glu	GTC Val	CCC Pro	ACC Thr	TAC Tyr 50	AAG Lys	TIG Leu	ATC Ile	TCC Ser	CAG Gln 55	TCT Ser	ATC Ile	TIG Leu	ATT Ile	GAC Asp 60		192
15	CGA Arg	CAC His	AAG Lys	GTT Val	AAC Asn 65	GGT Gly	TCC Ser	GTC Val	GCC Ala	CGA Arg 70	GCC Ala	GCT Ala	ATC Ile	CGA Arg	CAC His 75	CTT Leu		240
	GCC Ala	AAG Lys	GAG Glu	GGA Gly 80	TCC Ser	ATC Ile	AAG Lys	AAG Lys	ATT Ile 85	GTC Val	CAC His	CAC His	AAC Asn	GGA Gly 90	CAG Gln	Trp		288
20	ATC Ile	TAC Tyr	ACC Thr 95	CGA Arg	GCC Ala	ACT Thr	GCC Ala	GCT Ala 100	CCT Pro	GAC Asp	GCA Ala	TAAZ	ATCTO	EAT (		TEAT	G	341
25	GATY	TIG	AAA A	ATA	AAAA	A AA	YAAA!	AAA/	AA A									373
	(2)	INFO	ORMAI	MOI	FOR	SEQ	ID 1	<b>1</b> 0:43	3:									
30		,	(i) S	(B)	TYP	GTH: PE: a	: 103	RIST ami aci inea	imoa idi	cide	5							
35		i)	ii) N	10LEX	ULE	TYPE	E: pr	rotei	in									
		(2	ci) s	SEQUE	NŒ	DESC	RIP	MON:	SEX	) ID	NO:4	13:					•	
<b>4</b> 0	Met 1	Pro	Pro	Lys	Val 5	Lys	Ala	Lys	Thr	Gly 10	Val	Gly	Lys	Thr	Gln 15	Lys		
	Lys	Lys	Lys	Trp 20	Ser	Lys	Gly	Lys	Val 25	Lys	Asp	Lys	Ala	Ala 30	His	His		
45	Val	Val	Val 35	Asp	Gln	Ala	Thr	Tyr 40	Asp	Lys	Ile	Val	Lys 45	Glu	Val	Pro		
50	Thr	Tyr 50	Lys	Leu	Ile	Ser	Gln 55	Ser	Ile	Leu	Ile	60 Aap	Arg	His	Lys	Val		
	Asn 65	Gly	Ser	Val	Ala	Arg 70	Ala	Ala	Ile	Arg	His 75	Leu	Ala	Lys	Glu	Gly 80		
55	Ser	Ile	Lys	Lys	Ile 85	Val	His	His	Asn	Gly 90	Gln	Trp	Ile	Tyr	Thr 95	Arg		
	Ala	Thr	Ala	Ala 100	Pro	Asp	Ala											
60	(2)	DIFC	RMAI	MOI	FOR	Seq	ID 1	<b>1</b> 0:44	1:									
5.5		(i)	( <i>F</i> (E	QUENC A) LE B) T'S C) ST C) TC	NGII PE: RANI	i: 51 nucl	l4 ba Leic ESS:	ase p acid doub	pairs i	3								
		(ii)	MOI	ECUI	ΕT	Æ:	cDN7	Ā										

	(	iii)	HYP	OTHE	mc	L: N	Ö										
		(iv)	ANI	TI-SE	NSE:	NO											
5		(vi)	ORI ()			OURCE SM:		fia	rhod	юzyп	na						
10		(ix)	(E	A) N7 B) LC	ME/F	ŒY: ON: INFO	13		/pr	roduc	:t= "	'PRcI	INA78	3*1			
15	AAAA		SEC	TA TA	rs cn	TA T	C TO	T A	AA CZ	AG AA	AC AC	G AC					48
				IME:	1	:u 11	.e se	ar r?	75 GJ 5	in As	an Ai	g Ai		La I. LO	le Ph	æ	
20	GAG Glu	AAC Asn	CTC Leu 15	TTC Phe	AAG Lys	GAG Glu	GGA Gly	GTT Val 20	GCC Ala	GTC Val	GCC Ala	GCC Ala	AAG Lys 25	GAC Asp	TTC Phe	AAC Asn	96
25	GCT Ala	GCC Ala 30	ACC Thr	CAC His	CCC Pro	GAG Glu	ATT Ile 35	GAG Glu	GGT Gly	GTC Val	TCC Ser	AAC Asn 40	CIT Leu	GAG Glu	GTC Val	ATC Ile	144
30	AAG Lys 45	GCC Ala	ATG Met	CAG Gln	TCT Ser	TTG Leu 50	ACC Thr	TCC Ser	AAG Lys	GGA Gly	TAC Tyr 55	GTG Val	AAG Lys	ACC Thr	CAG Gln	TTC Phe 60	192
35	TCG Ser	TGG Trp	CAG Gln	TAC Tyr	TAC Tyr 65	TAC Tyr	TAC Tyr	ACC Thir	CIC Leu	ACC Thr 70	CCT Pro	GAG Glu	GGT Gly	CIT Leu	GAC Asp 75	TAC Tyr	240
33	CTC Leu																288
40	AAG Lys	CGA Arg	CCC Pro 95	ACC Thr	CGA Arg	CCT Pro	GCC Ala	AAG Lys 100	GCC Ala	CAG Gln	GGT Gly	CCC Pro	GGA Gly 105	GGT Gly	GCC Ala	TAC Tyr	336
45	CGA Arg	GCT Ala 110	Pro	CGA Arg	GCC Ala	GAG Glu	GGT Gly 115	GCC Ala	GGT Gly	CGA Arg	GGA Gly	GAG Glu 120	TAC Tyr	CGA Arg	CGA Arg	CGA Arg	384
50	GAG Glu 125	GAC Asp	Gly	GCC Ala	GGT Gly	GCC Ala 130	Phe	GGT Gly	GCC	Gly	CGA Arg 135	Gly	GGA	. CCC Pro	CGA Arg	GCT Ala 140	432
	TAA	ATCC	CAG.	AGCI	TTTC	TT T	TIGI	CCIT	G CI	GGGA	CTAT	GGC	ATGA	TGA	GCTG	GCTTG(	2 492
55	AGA	AAAA	AAA.	AAAA	AAAA	AA A	A										514
	(2)	INF	ORMA	TION	FOR	SEQ	ID	NO:4	5:								
60			(i)	() (E	l) LE 3) TY	CHA NGIH PE: POLC	: 14 amin	ο aπ	ino id		ls						
65		•	(ii)	MOLE	CULE	TYF	E: p	rote	in								
		1	(xi)	SEQU	ENCE	DES	CRIE	TIO	1: SI	Ωп	) NO:	45:					
	Met 1		ı Ile	e Ser	Lys		) Ası	ı Arç	J Arg	g Ala 10		e Phe	e Glı	ı Ası	n Let	Phe	

	Lys G	lu G	ly Val 20	Ala	Val	Ala	Ala	Lys 25	Asp	Phe	: Asn	Ala	Ala 30		His	
5	Pro G	lu I	le Glu 35	Gly	Val	Ser	Asn 40	Leu	Glu	Val	Ile	Lys 45	Ala	Met	Gln	
	Ser L	eu 11 50	hr Ser	. TÀ2	Gly	Tyr 55	Val	Lys	Thr	Gln	Phe 60	Ser	Trp	Gln	Tyr	
10	Tyr T 65	yr T	yr Thr	Leu	Thr 70	Pro	Glu	Gly	Leu	Asp 75	Tyr	Leu	Arg	Glu	Phe 80	
15	Leu H	is Le	eu Pro	Ser 85	Glu	Ile	Val	Pro	Asn 90	Thr	Leu	Lys	Arg	Pro 95		
	Arg P	ro Al	la Lys 100	Ala	Gln	Gly	Pro	Gly 105	Gly	Ala	Tyr	Arg	Ala 110	Pro	Arg	
20	Ala G	lu Gl	ly Ala 15	Gly	Arg	Gly	Glu 120	Tyr	Arg	Arg	Arg	Glu 125	Asp	Gly	Ala	
	Gly Al	la Ph 30	æ Gly	Ala	Gly	Ar <del>g</del> 135	Gly	Gly	Pro	Arg	Ala 140					
25	(2) <b>n</b>	VFORM	ation	FOR	SEQ	ID N	<b>1</b> 0:46	5:								
30	ı	(i) S	EQUENT (A) LI (B) T (C) S (D) T	ENGTI YPE : IRANI	i: 43 nucl	87 ba Leic ESS:	se p acid doub	airs l	S							
35	(i	.i) M	DLECU													
	(ii	.i) H	YPOTHE	TIC	AL: N	Ö				-						
	i)	.v) A	NTI-SI	NSE:	NO											
40	(√	ri) O	RIGINA (A) OF	AL SC SCANI	JURCE SM:	: Phaf	fia	rhod	lozym	a						
45	, <b>(i</b>		EATURE (A) NA (B) LC (D) OI	ME/K	ON:	30	308 ION:	/pr	oduc	t= "	PRcD	<b>NA8</b> 5	n			
50	(x	i) s	EQUENC	E DE	SCRI	PIIO	N: S	EQ I	סאו ס	:46:						
	CICCCI	CAAG	YTAAA	AACC	A CC	GCAC	ATC .	ATG Met 1	TCC : Ser :	AAG Lys .	CGA . Arg '	ACC I	AAG : Lys :	aaa Lys	GTT Val	53
55	GGA AT Gly Il	- 11H	C GGA r Gly	AAG Lys	TAC Tyr	GGA ( Gly ) 15	GIC ( Val )	CGA Arg	TAC ( Tyr (	3GA ( Gly )	GCT   Ala   20	rcc ( Ser 1	Erc (	OGA Arg	AAG Lys	101
60	ACC GT Thr Va 25	C AA 1 Ly:	G AAG s Lys	NTG ( Xaa (	GAG Glu 30	GIC '	TCG (	CAG ( Gln )	CAC ( His (	GT : Sly ' 35	ACC '	TAC I	ACC :	IGT Cys	GAC Asp 40	149
65	TTC TG Phe Cy	C GG S Gly	A AAG y Lys	GAC Asp 45	GCC ( Ala '	GIC I	AAG ( Lys )	OGA / Arg '	ACC (Thr )	GCT ( Ala '	GTT ( Val (	GT A	ATC :	rec Imp 55	AAG Lys	197
70	TGC CG Cys An	A GGZ g Gly	A TGC / Cys 60	CGA / Arg :	AAG : Lys '	ACC I	ACC ( Ihr i	GCC ( Ala ( 65	Gly (	er (	GCT : Ala :	rp (	AG ( Eln 1	err Leu	CAG Gln	245

	ACC ACC GCC GCT CTC ACC GTC AAG TCC ACC ACT CGA CGA CTC CGA GAG Thr Thr Ala Ala Leu Thr Val Lys Ser Thr Thr Arg Arg Leu Arg Glu 75 80 85	293
s	CTC AAG GAG GIT TAAATTGAAT TCTGCACAAA GACAAAACTG TTGCGGGGGG Leu Lys Glu Val 90	345
_	GAGAGAGTOG ATTCATTCTT TTTTTTTGTA GATCTGAAGG GATGCCATGT CAACCCTTTC	405
0	GITCCCCAAA AAAAAAAAA AAAAAAAAA AA	437
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	<ul><li>(i) SEQUENCE CHARACTERISTICS:</li><li>(A) LENGTH: 92 amino acids</li><li>(B) TYPE: amino acid</li><li>(D) TOPOLOGY: linear</li></ul>	
TC	(ii) MOLECULE TYPE: protein	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:47:	
<u>.</u> ,	Met Ser Lys Arg Thr Lys Lys Val Gly Ile Thr Gly Lys Tyr Gly Val 1 5 10 15	
	Arg Tyr Gly Ala Ser Leu Arg Lys Thr Val Lys Lys Xaa Glu Val Trp 20 25 30	
<b>X</b> O	Gln His Gly Thr Tyr Thr Cys Asp Phe Cys Gly Lys Asp Ala Val Lys 35 40 45	
35	Arg Thr Ala Val Gly Ile Trp Lys Cys Arg Gly Cys Arg Lys Thr Thr 50 55 60	
	Ala Gly Gly Ala Trp Gln Leu Gln Thr Thr Ala Ala Leu Thr Val Lys 65 70 75 80	
<b>4</b> 0	Ser Thr Thr Arg Arg Leu Arg Glu Leu Lys Glu Val 85 90	
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<b>5</b> 0	(C) STRANDEDNESS: double (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: CENA	
55	(iii) HYPOTHETICAL: NO	
	(iv) ANTI-SENSE: NO	
	(vi) ORIGINAL SOURCE:  (A) ORGANISM: Phaffia rhodozyma	
60 دئ	(ix) FEATURE: (A) NAME/KEY: CDS (B) LOCATION: 35400 (D) OTHER INFORMATION: /product= "PRoDNA87"	
w	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:48:	
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70	Met Ala Thr Lys Thr Gly	

												1				5		
5	AA Ly:	G AC	r	CGA Arg	Ser 10	WIG	CTC Leu	CAG Gln	GAC Asp	GIC Val 15	Val	' ACT Thr	Arg CGG	GAG Glu	TAC Tyr 20	Thr	ATC	100
10	CA:	C CI	C u	CAC His 25	AAG Lys	TAC	GTT Val	CAC His	GGA Gly 30	Arg	TCT	TTC Phe	AAG Lys	AAG Lys 35	CGA Arg	GCT Ala	CCT	148
	TG		T a 0	GTC Val	AAG Lys	TCC	ATC	CAG Gln 45	GAG Glu	TTT Phe	GCT Ala	CTC Leu	AAG Lys 50	TCG Ser	ATG Met	GGA Gly	ACC Thr	196
15																		
20	Arc 55	, ~	T P	GIC Val	CGA Arg	ATT Ile	GAC Asp 60	CCC Pro	AAG Lys	TIG Leu	AAC Asn	CAG Gln 65	GCÇ Ala	GTC Val	TGG Trp	GGA Gly	CAG Gln 70	244
	Gly	' GT ' Va	C ;	AAG Lys	AAC Asn	CCC Pro 75	CCC Pro	AAG Lys	CGA Arg	CTC Leu	CGA Arg 80	ATC Ile	CGA Arg	CTT Leu	GAG Glu	CGA Arg 85	AAG Lys	292
25	CGA Arg	AA As	C ( n ;	GAC Asp	GAG Glu 90	GAG Glu	gat Asp	GCT Ala	AAG Lys	GAC Asp 95	AAG Lys	CTC Leu	TAC Tyr	ACT Thr	CTT Leu 100	GCT Ala	ACC Thr	340
30	GTC Val	Va.		200 Pro 105	GGA Gly	GTC Val	ACC Thr	AAC Asn	TTC Phe 110	AAG Lys	GGT Gly	CTC Leu	Gln	ACC Thr 115	GTT Val	GTC Val	GTT Val	388
35	GAC Asp	Thi 120		SAG Slu	<b>Laa</b> t	TTTC	arc 1	TGGA	TTT	TA O	GACG	GICG	ATT	CAGC	rgr			437
	TIC	TIG	300	3C C	ATTC	TICI	TA T	GCAC	ICIG	ATG	CCTI	TCA	CGAC	CONI	TT T	INIT	TCINA	497
40				A A														509
	(2)	IN	7OF	MAT	IQN	FOR	SEQ	ID N	0:49									
45					EQUE (A) (B)	NCE LEN TYP	CHAR GIH: E: a	ACTE 121 mino Y: 1	RIST ami	ICS: no a d	cids							
		(	ii	.) M	OLEC	ULE	TYPE	: pr	otei	n								
50		(	xi	.) s	EQUE	NCE	DESC	RIPT	ION:	SEQ	ID :	NO:4:	9:					
**	Met 1											Ala 1		Sln i	Asp 1	/al ' 15	Val	
55	Thr	Arg	G	lu 1	Iyr ' 20	Thr	Ile :	His 1	Leu 1	His 1 25	Lys '	Tyr v	Val I	lis (	Gly <i>1</i> 30	Arg:	Ser	
60	Phe	Lys	L	ys 2 35	Arg i	Ala :	Pro '	Trp i	Ala 1 40	Val 1	Lys :	Ser :	Ile (	Sln ( 45	Glu I	Phe 1	Ala	
	Leu	Lys 50	S	er 1	Met (	Gly '	Thr i	Arg i	Asp 1	Val 2	Arg :	Ile A	Asp I 60	Pro I	Lys I	.eu 1	Asn	
65	Gln 65	Ala	V	al :	) qrí	Gly (	Gln ( 70	Gly V	/al 1	Lys 1	Asn 1	Pro I 75	Pro I	ys 1	Arg I	.eu )	80 Arg	
70	Ile	Arg	L	eu (	Glu A	Arg 1 85	Lys 1	Arg A	Asn A	Asp (	3lu ( 90	Glu A	Asp A	la I	Jys A	Asp I 95	ràz	

	Leu Tyr Thr Leu Ala Thr Val Val Pro Gly Val Thr Asn Phe Lys Gly 100 105 110	
5	Leu Gln Thr Val Val Val Asp Thr Glu 115 120	
	(2) INFORMATION FOR SEQ ID NO:50:	
10	(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 542 base pairs  (B) TYPE: nucleic acid	
15	(C) STRANDEINESS: double (D) TOPOLOGY: linear (ii) MOLECULE TYPE: CDNA	
	(iii) HYPOTHETICAL: NO	
20	(iv) ANTI-SENSE: NO	
	(vi) ORIGINAL SOURCE: (A) ORGANISM: Phaffia rhodozyma	
프	(ix) FEATURE:  (A) NAME/KEY: CDS  (B) LOCATION: 18443  (D) OTHER INFORMATION: /product= "PRoDNA95"	
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35	AGTOGOTATA CATORAG ATG TOO GTO GOT GTO CAG ACT TTO GGT AAG AAG Met Ser Val Ala Val Gln Thr Phe Gly Lys Lys 1 5 10	50
	AAG ACT GCC ACC GCT GTG GCC CAC GCC ACC CCT GGC CGA GGT CTC ATC Lys Thr Ala Thr Ala Val Ala His Ala Thr Pro Gly Arg Gly Leu Ile 15 20 25	98
40	CGA CTT AAC GGA CAG CCT ATC TCA CTT GCC GAG CCT GCT CTC CTC CGA Arg Leu Asn Gly Gln Pro Ile Ser Leu Ala Glu Pro Ala Leu Leu Arg 30 35 40	146
45	TAC AAG TAC TAC GAG CCT ATC CTC GTC ATC GGA GCT GAG AAG ATC AAC Tyr Lys Tyr Tyr Glu Pro Ile Leu Val Ile Gly Ala Glu Lys Ile Asn 45 50 55	194
50	CAG ATC GAC ATC CGA CTC AAG GTC AAG GGT GGA GGA CAC GTC TCC CAG Gln Ile Asp Ile Arg Leu Lys Val Lys Gly Gly Gly His Val Ser Gln 60 65 70 75	242
55	GTG TAC GCC GTC CGA CAG GCC ATC GGT AAG GCC ATC GTC GCT TAC TAC Val Tyr Ala Val Arg Gln Ala Ile Gly Lys Ala Ile Val Ala Tyr Tyr 80 85 90	290
	GCT AAG AAC GTC GAT GCC GCC TCT GCC CTC GAG ATC AAG AAG GCT CTC Ala Lys Asn Val Asp Ala Ala Ser Ala Leu Glu Ile Lys Lys Ala Leu 95 100 105	338
60	GTC GCC TAC GAC CGA ACC CTC CTC ATC GCC GAT CCC CGA CGA ATG GAG Val Ala Tyr Asp Arg Thr Leu Leu Ile Ala Asp Pro Arg Arg Met Glu 110 115 120	386
65	CCC AAG AAG TTC GGA GGA CCC GGA GCC CGA GCC CGA GTC CAG AAG TCT Pro Lys Lys Phe Gly Gly Pro Gly Ala Arg Ala Arg Val Gln Lys Ser 125 130 135	434
7	TAC CGA TAAAAAGIGI TIGICITGIG GICIGGOGG TCATCIATCC AACATCITIG  Tyr Arg	490

# GAAAANANIT GITTOGGICA TATGICATGC CTCTTTATOG AAAAAAAAAA AA

542

- (2) INFORMATION FOR SEQ ID NO:51:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGIH: 141 amino acids
    - (B) TYPE: amino acid
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:51:

Met Ser Val Ala Val Gln Thr Phe Gly Lys Lys Lys Thr Ala Thr Ala 1 5 10 15

20 Val Ala His Ala Thr Pro Gly Arg Gly Leu Ile Arg Leu Asn Gly Gln 20 25 30

Pro Ile Ser Leu Ala Glu Pro Ala Leu Leu Arg Tyr Lys Tyr Tyr Glu 35 40 45

Pro Ile Leu Val Ile Gly Ala Glu Lys Ile Asn Gln Ile Asp Ile Arg 50 55 60

Leu Lys Val Lys Gly Gly Gly His Val Ser Gln Val Tyr Ala Val Arg 55 70 75 80

Gln Ala Ile Gly Lys Ala Ile Val Ala Tyr Tyr Ala Lys Asn Val Asp 85 90 95

35 Ala Ala Ser Ala Leu Glu Ile Lys Lys Ala Leu Val Ala Tyr Asp Arg 100 105 110

Thr Leu Leu Ile Ala Asp Pro Arg Arg Met Glu Pro Lys Lys Phe Gly
115 120 125

Gly Pro Gly Ala Arg Ala Arg Val Gln Lys Ser Tyr Arg 130 135

#### Claims

- 1. Recombinant DNA comprising a transcription promoter and a downstream sequence to be expressed, in operable linkage therewith,
- wherein the transcription promoter comprises a region found upstream of the open reading frame of a highly expressed *Phaffia* gene.
- 2. Recombinant DNA according to claim 1, wherein said highly expressed *Phaffia* gene is a glycolytic pathway gene.
- 3. Recombinant DNA according to claim 2, wherein said glycolytic pathway gene is a gene coding for Glyceraldehyde-3-Phosphate Dehydrogenase.
- 4. Recombinant DNA according to claim 1, wherein said highly expressed *Phaffia* gene is a ribosomal protein encoding gene.
  - 5. Recombinant DNA comprising a transcription promoter and a downstream sequence to be expressed, in operable linkage therewith,

wherein the transcription promoter comprises a region found upstream of the open reading frame encoding a protein as represented by one of the amino acid sequences depicted in any one of SEQIDNOs: 24 to 50.

- 6. A recombinant DNA according to any one of the preceding claims, wherein said downstream sequence to be expressed is heterologous with respect to the transcription promoter sequence.
- A recombinant DNA according to any one of claims 1 to 6, wherein the downstream sequence comprises an open reading frame coding for a polypeptide responsible for reduced sensitivity against a selective agent.
- 30 8. A recombinant DNA according to claim 7, wherein said selective agent is G418.
  - 9. A recombinant DNA according to any one of claims 1 to 6, wherein the said downstream sequence to be expressed codes for an enzyme involved in the carotenoid biosynthesis pathway.
- 10. A recombinant DNA according to claim 9, wherein said downstream sequence to be expressed encodes an enzyme having an activity selected from the group consisting of isopentenyl pyrophosphate isomerase, geranylgeranyl pyrophosphate synthase, phytoene synthase, phytoene desaturase, and lycopene cyclase.

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- 11. A recombinant DNA according to claim 10, wherein said downstream sequence to be expressed encodes an enzyme having an amino acid sequence selected from the one represented by SEQIDNO: 13, SEQIDNO: 15, SEQIDNO: 17, SEQIDNO: 19, SEQIDNO: 21 or SEQIDNO: 23.
- 12. A recombinant DNA according to any one of the preceding claims, wherein said recombinant DNA comprises further a transcription terminator downstream from the said DNA sequence to be expressed, in operable linkage therewith.
- 13. A recombinant DNA according to claim 12, wherein the terminator is a GAPDH-encoding gene terminator fragment.
  - 14. A recombinant DNA according to any one of the preceding claims, wherein the recombinant DNA is in the form of a vector capable of replication and/or integration in a host organism.
- 15. A recombinant DNA according to claim 14, further comprising *Phaffia* ribosomal RNA encoding DNA.
  - 16. A recombinant DNA according to claim 15, which is linearised by cleaving inside the *Phaffia* ribosomal RNA encoding DNA portion.
  - 17. A microorganism harbouring a recombinant DNA according to any one of the preceding claims.
  - 18. A microorganism according to claim 17, which is Phaffia rhodozyma.
- 25 19. A microorganism according to claim 18, having the recombinant DNA integrated into its genome in an amount of 50 copies or more.
  - 20. An isolated DNA fragment comprising a *Phaffia* GAPDH-gene, or a functional fragment thereof.
  - 21. Use of a functional fragment according to claim 20 for making a recombinant DNA construct.
  - 22. The use according to claim 21, wherein said fragment is a regulatory region normally located upstream or downstream of the open reading frame coding for GAPDH in *Phaffia rhodozyma*.
  - 23. A method for obtaining a transformed *Phaffia* strain, comprising the steps of
    (a) contacting cells or protoplasts of a *Phaffia* strain with recombinant DNA under conditions conducive to uptake thereof,

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said recombinant DNA comprising a transcription promoter and a downstream sequence to be expressed in operable linkage therewith,

- (b) identifying *Phaffia rhodozyma* cells or protoplasts having obtained the said recombinant DNA in expressible form,
  - wherein the recombinant DNA is one according to any one of the preceding claims.
- A method according to claim 23, comprising the additional step of providing an electropulse after contacting of *Phaffia* cells or protoplasts with the said recombinant DNA.
- 25. A transformed *Phaffia* strain obtainable by a method according to any one of the preceding claims, said strain, upon cultivation, being capable of expression of the said downstream sequence, as a consequence of transformation with the said recombinant DNA.
- 26. A transformed *Phaffia* strain according to claim 25, wherein the said downstream sequence codes for a pharmaceutical protein.
  - 27. A transformed *Phaffia* strain according to any one of claims 24 to 26, wherein the said *Phaffia* strain contains at least 10, preferably at least 50, copies of the said recombinant DNA integrated into its genome.
  - 28. An isolated DNA sequence coding for an enzyme involved in the carotenoid biosynthetic pathway of *Phaffia rhodozyma*.
- 29. An isolated DNA sequence according to claim 28, wherein said enzyme has an activity selected from isopentenyl pyrophosphate isomerase activity, geranylgeranyl pyrophosphate synthase activity, phytoene synthase activity, phytoene desaturase activity and lycopene cyclase activity.
  - 30. An isolated DNA sequence coding for an enzyme having an amino acid sequence selected from the one represented by SEQIDNO: 13, SEQIDNO: 15, SEQIDNO: 17, SEQIDNO: 19, SEQIDNO: 21 or SEQIDNO: 23.
    - 31. An isolated DNA sequence coding for a variant of an enzyme according to claim 30, said variant being selected from (i) an allelic variant, (ii) an enzyme having one or more amino acid additions, deletions and/or substitutions and still having the stated enzymatic activity.
    - 32. An isolated DNA sequence encoding an enzyme involved in the carotenoid biosynthesis pathway selected from:
    - (i) a DNA sequence as represented in SEQIDNO: 12, SEQIDNO: 14, SEQIDNO: 16 SEQIDNO: 18; SEQIDNO: 20, or SEQIDNO: 22,

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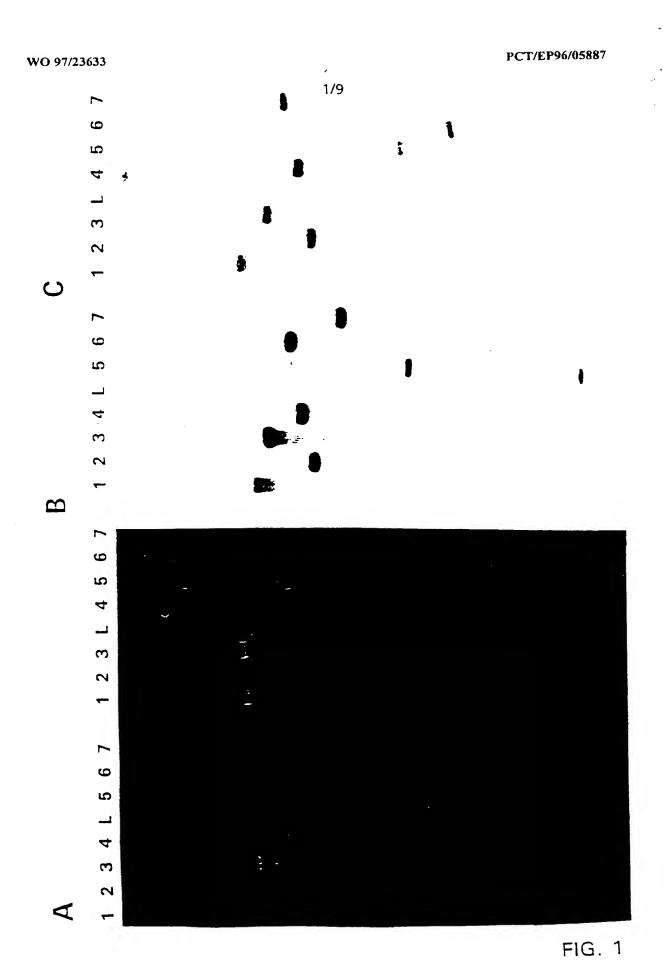
- (ii) an isocoding variant of the DNA sequence represented in SEQIDNO: 12, SEQIDNO: 14, SEQIDNO: 16, SEQIDNO: 18, SEQIDNO: 20 or SEQIDNO: 22;
- (iii) an allelic variant of a DNA sequence as represented in SEQIDNO: 12, SEQIDNO: 14, SEQIDNO: 16, SEQIDNO: 18; SEQIDNO: 20 or SEQIDNO: 22;
- (iv) a DNA sequence capable, when bound to nitrocellulose filter and after incubation under hybridising conditions and subsequent washing, of specifically hybridising to a radio-labelled DNA fragment having the sequence represented in SEQIDNO: 12, SEQIDNO: 14, SEQIDNO: 16, SEQIDNO: 18, SEQIDNO: 20 or SEQIDNO: 22, as detectable by autoradiography of the filter after incubation and washing, wherein said incubation under hybridising conditions and subsequent washing is performed by incubating the filter-bound DNA at a temperature of at least 50°C, preferably at least 55°C, in the presence of a solution of the said radio-labeled DNA in 0.3 M NaCl, 40 mM Tris-HCl, 2 mM EDTA, 0.1% SDS, pH 7.8 for at least one hour, whereafter the filter is washed at least twice for about 20 minutes in 0.3 M NaCl, 40 mM Tris-HCl, 2 mM EDTA, 0.1% SDS, pH 7.8, at a temperature of 50°C, preferably at least 55°C, prior to autoradiography.
  - 33. Recombinant DNA comprising an isolated DNA sequence according to any one of claims 27 to 32.
- 34. Recombinant DNA according to claim 33, wherein said isolated DNA sequence is operably linked to a transcription promoter capable of being expressed in a suitable host, said isolated DNA sequence optionally being linked also to a transcription terminator functional in the said host.
  - 35. Recombinant DNA according to claim 34, wherein said host is a *Phoffia* strain.
- 25 36. Recombinant DNA according to any one of claims 33 to 35, wherein the transcription promoter is from a glycolytic pathway gene present in *Phaffia*.
  - 37. Recombinant DNA according to claim 36, wherein said glycolytic pathway gene is a gene coding for Glyceraldehyde-3-Phosphate Dehydrogenase.
  - 38. Recombinant DNA according to any one of claims 33 to 35, wherein the transcription promoter is from a ribosomal protein encoding gene.
  - 39. Recombinant DNA according to any one of claims 33 to 35, wherein the transcription promoter comprises a region found upstream of the open reading frame encoding a protein as represented by one of the amino acid sequences depicted in any one of SEQIDNOs: 24 to 50.

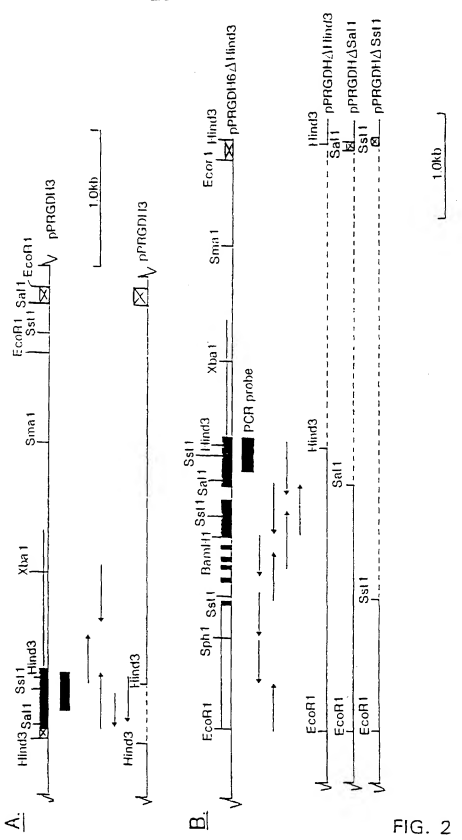
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- 40. Recombinant DNA according to any one of claims 27 to 39, wherein said recombinant DNA comprises further a transcription terminator downstream from the said heterologous DNA sequence to be expressed, in operable linkage therewith, which terminator is a *Phaffia* transcription terminator.
- 41. Recombinant DNA according to any one of claims 27 to 40, being in the form of a vector.
  - 42. Use of a vector according to claim 41 to transform a host.
  - 43. Use according to claim 19, wherein the host is a *Phaffia* strain.
  - 44. A host obtainable by transformation, optionally of an ancestor, using a recombinant DNA according to any one of claims 27 to 41.
  - 45. A host according to claim 44, which is a Phaffia strain, preferably a Phaffia rhodozyma strain.
  - 46. A transformed *Phaffia rhodozyma* strain which is capable of overexpressing a DNA sequence encoding an enzyme involved in the carotenoid biosynthesis pathway.
- 47. A transformed *Phaffia rhodozyma* strain according to claim 46, which produces inreased amounts of astaxanthin relative to its untransformed ancestor.
  - 48. A method for producing an enzyme involved in the carotenoid biosynthesis pathway, by culturing a host according to claim 44 or 45, under conditions conducive to the production of said enzyme.
  - 49. A method for producing a carotenoid, characterised in that a host according to any one of claims 44 to 47 is cultivated under conditions conducive to the production of the carotenoid.
  - 50. A method according to claim 49, wherein the carotenoid is astaxanthin.
  - 51. A method for producing a pharmaceutical protein by culturing a transformed *Phaffia* strain according to claim 26 under conditions conducive to the production of the said protein.
  - 52. A method for the isolation of a promoter from a highly expressed gene in *Phaffia*, comprising the steps of:
    - (a) making a cDNA library on mRNA isolated from a Phaffia strain grown under desired conditions;
    - (b) determining (part of) the nucleotide sequence of the (partial) cDNAs obtained in step (a);
    - (c) comparing the obtained sequence data in step (b) to known sequence data;

- (d) cloning amplifying putative promoter fragments of the gene located either directly upstream of the open reading frame or directly upstream of the transcription start site of the gene corresponding to the expressed cDNA, and
- (e) verifying whether the promoter sequences obtained give high-level expression in a *Phaffia* strain, by expressing a suitable marker under the control of the promoter in a transformed *Phaffia* strain.





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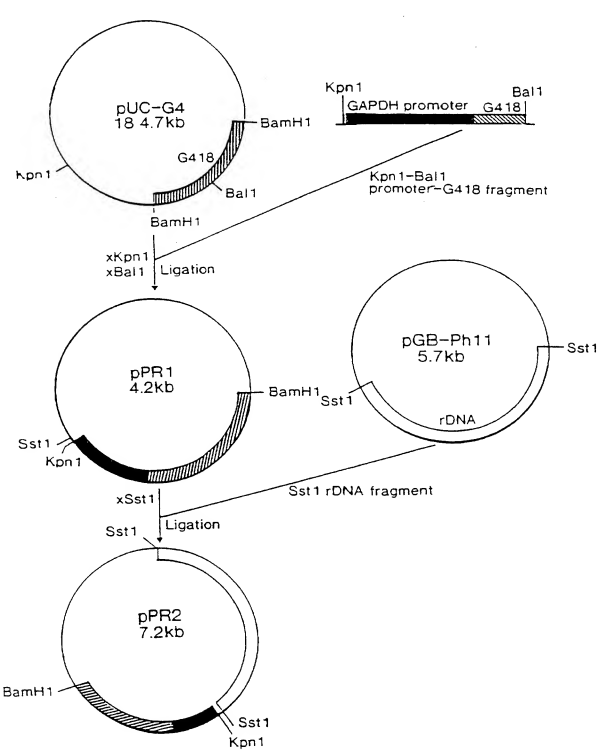


FIG. 3

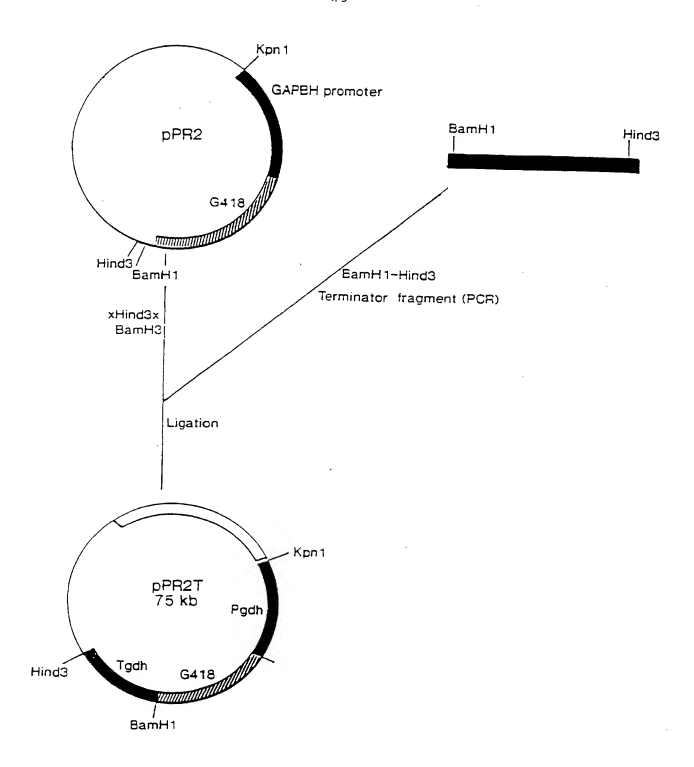


FIG. 4

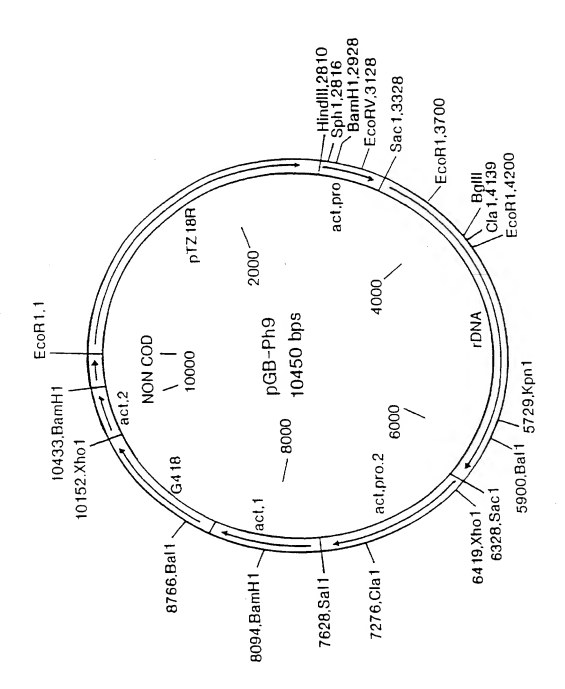
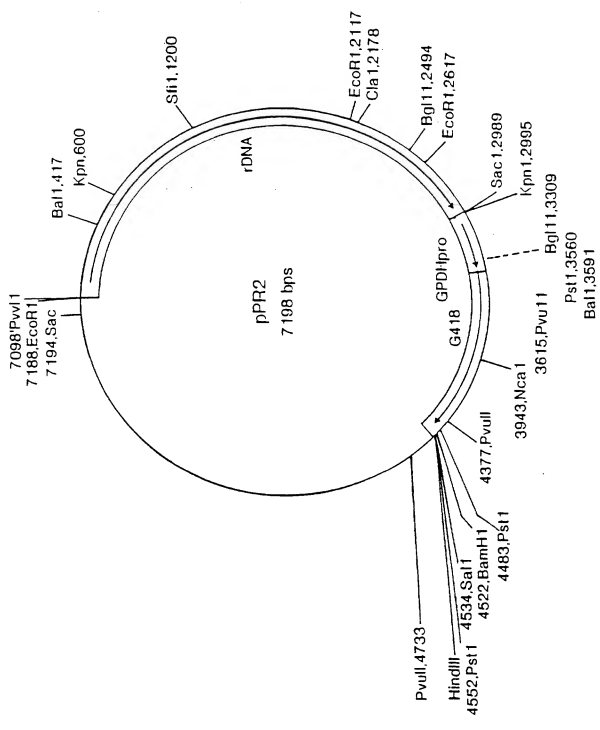


FIG. 5



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FIG. 6

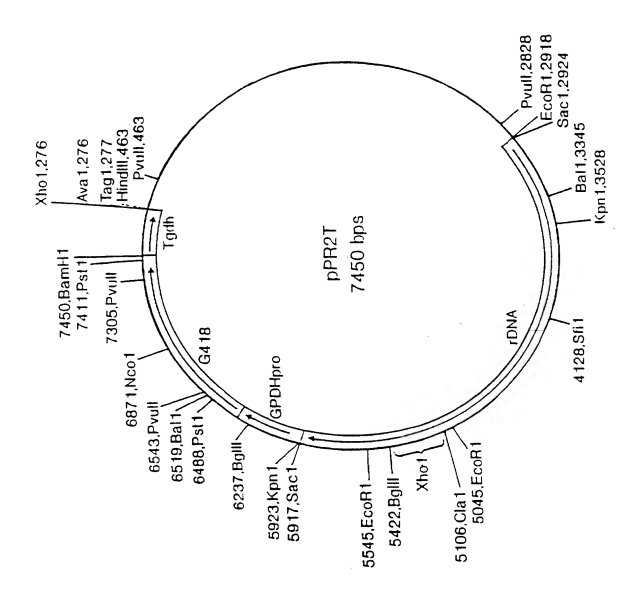


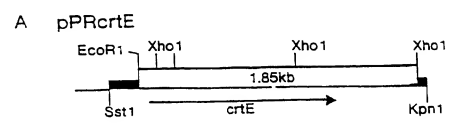
FIG. 7

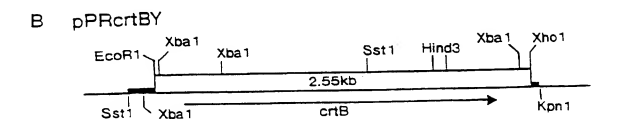
# Carotenoid Biosynthetic Pathway of Erwinia uredovora

Farnesyl Pyrophosphate (FPP) + Isopetenyl Pyrophosphate (GPP)

FIG. 8

Zeaxanthin-Diglucoside





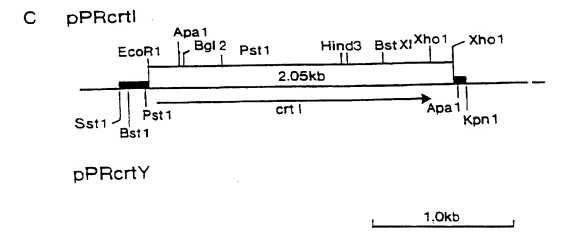


FIG. 9

Internation 'Application No PCT/EP 96/05887

			PCT/LP 96/05887
IPC 6	C12N15/81 C12N1/16 C07K14 C12N15/52 C12N15/60 C12P23 C12R1:645),(C12N1/21,C12R1:19)	/00 C12N1/2	2 C12N15/53 1 //(C12N1/16,
	to International Patent Classification (IPC) or to both national cla S SEARCHED	assification and IPC	
	S SEARCHED  documentation searched (classification system followed by classifi	cation symbols)	
IPC 6	C12N C07K C12P	cauch symmosy	
Documenta	tion searched other than minimum documentation to the extent th	at such documents are inclu	uded in the fields searched
Electronic	data base consulted during the international search (name of data	base and, where practical, s	search terms used)
C. DOCUM	MENTS CONSIDERED TO BE RELEVANT		
Category *	Citation of document, with indication, where appropriate, of the	e relevant passages	Relevant to claim No.
X	ANALES DE LA REAL ACADEMIA DE Fo vol. 61, no. 4, 1995, pages 463-471, XP000577134 J. ANDRIO ET AL.: "Transformac Phaffia rhodozyma utilizando el acetato de litio."	iôn de	1,6-8, 12,14, 17-19, 23,25, 27, 33-35, 40,44,45
	summary, page 463, page 468, pa see page 464, paragraph 1	ragraph 3	10,11,13
X	EP 0 590 707 A (GIST BROCADES N 1994 cited in the application	V) 6 April	1,6-12, 14, 17-19, 23-25, 27-35,
Y	see the whole document		40-50 26,51,52
		-/	
X Furt	ther documents are listed in the continuation of box C.	X Patent family m	nembers are listed in annex.
"A" docum	stegones of cited documents :	"T" later document publi	ished after the international filing date I not in conflict with the application but the principle or theory underlying the
"E" earlier filing	tered to be of particular relevance document but published on or after the international date ent which may throw doubts on priority claim(s) or	"X" document of particu cannot be considere	ular relevance; the claimed invention ed novel or cannot be considered to
which citatio	is cited to establish the publication date of another in or other special reason (as specified)  nent referring to an oral disclosure, use, exhibition or means	"Y" document of particu- cannot be considere document is combine	e step when the document is taken alone  alar relevance; the claimed invention  ed to involve an inventive step when the  ned with one or more other such docu-  tation being obvious to a person skilled
iater t	ent published prior to the international filing date but han the priority date claimed	in the art.	of the same patent family
	actual completion of the international search  June 1997	-	ne international search report 1.2.06.97
Name and	mailing address of the ISA  European Patent Office, P.B. 5818 Patentiaan 2  NL - 2280 HV Rijswijk  Tel. (+ 31-70) 340-2040, Tx. 31 651 epo nl,  Fax: (+ 31-70) 340-3016	Authorized officer	

Form PCT/ISA/218 (second sheet) (July 1992)

Internation 'Application No PCT/EP 96/05887

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Form PCT/ISA/210 (continuation of second sheet) (July 1992)

Interna* d Application No
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Internati 'Application No

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Form PCT/ISA/210 (continuation of second sheet) (July 1992)

Inti Lional application No.

PCT/EP 96/05887

Box I Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)
This International Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:
Claims Nos.:     because they relate to parts of the international Application that do not comply with the prescribed requirements to such an extent that no meaningful international Search can be carried out, specifically:
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Classes Nos.:  because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
Bex II Observations where unity of invention is lacking (Continuation of item 2 of first sheet)
This International Searching Authority found multiple inventions in this international application, as follows:
see continuation-sheet
As all required additional search fees were timely paid by the applicant, this International Search Report covers all searchable claims.
2. X As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
3. As only some of the required additional search fees were timely paid by the applicant, this International Search Report covers only those claims for which fees were paid, specifically claims Nos.:
4. No required additional search fees were timely paid by the applicant. Consequently, this International Search Report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:
Remark on Protest  The additional search fees were accompanied by the applicant's protest.  No protest accompanied the payment of additional search fees.

Form PCT/ISA/210 (continuation of first sheet (1)) (July 1992)

International Application No. PCT/EP 96/05887

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### FURTHER INFORMATI N CONTINUED FROM PCT/ISA/210

- 1. Recombinant DNA comprising a transcription promoter and downstream region to be expressed where the transcription promoter comprises a region found upstream of a highly expressed Phaffia gene, method of transforming a Phaffia strain where the transcription promoter is from a glycolytic pathway gene, to express a downstream sequence, recombinant DNA thereof, including a selective agent and the transformed Phaffia strains: Claims 2, 3, 13, 36 and 37 (completely) and Claims 1, 6 to 14, 17 to 19, 22 to 27, 33 to 35 and 40 to 45 and 51 (partially).
- 2. Recombinant DNA comprising a transcription promoter and downstream region to be expressed where the transcription promoter comprises a region found upstream of a highly expressed Phaffia gene, method of transforming a Phaffia strain where the transcription promoter is from a ribosomal protein, to express a downstream sequence, recombinant DNA thereof and the transformed Phaffia strains: Claims 4, 5, 15, 16, 38 and 39 (completely) and Claims 1, 6 to 12, 14, 17 to 19, 22 to 27, 33 to 35 and 40 to 45 and 51 (partially).
- 3. An isolated DNA fragment comprising a <u>Phaffia GAPDH-gene</u> and use in the construction of a DNA construct: <u>Claims 20 to 21</u> {completely} and <u>Claim 22</u> {partially}.
- 4. An isolated DNA sequence coding for an enzyme involved in the carotenoid biosynthetic pathway of Phaffia rhodozyme and recombinant DNA comprising a transcription promoter and the downstream region to be expressed codes for an enzyme involved in the carotenoid biosynthesis pathway and the transformed Phaffia strains comprising said DNA: Claims 1.6. 9 to 12.14.17 to 19. 23 to 27. 28 to 35 and 40 to 50 {partially}
- 5. An isolated DNA sequence coding for an enzyme involved in the carotenoid biosynthetic pathway of Phaffia rhodozyme, and recombinant DNA comprising a transcription promoter and the downstream region to be expressed codes for an enzyme involved in the carotenoid biosynthesis pathway and the transformed Phaffia strains comprising said DNA, where the enzyme has **isop ntenyl pyrophosphate isomerase activity**: Claims 1, 6, 9 to 12, 14, 17 to 19, 23 to 27, 28 to 35 and 40 to 50 {partially}

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#### FURTHER INFORMATION CONTINUED FROM PCT/ISA/210

- 6. An isolated DNA sequence coding for an enzyme involved in the carotenoid biosynthetic pathway of Phaffia rhodozyme, and recombinant DNA comprising a transcription promoter and the downstream region to be expressed codes for an enzyme involved in the carotenoid biosynthesis pathway and the transformed Phaffia strains comprising said DNA, where the enzyme has **geranylgeranyl** pyrophosphate synthase activity: Claims 1, 6, 9 to 12, 14, 17 to 19, 23 to 27, 28 to 35 and 40 to 50 {partially}
- 7. An isolated DNA sequence coding for an enzyme involved in the carotenoid biosynthetic pathway of Phaffia rhodozyme and recombinant DNA comprising a transcription promoter and the downstream region to be expressed codes for an enzyme involved in the carotenoid biosynthesis pathway and the transformed Phaffia strains comprising said DNA, where the enzyme has phytoene sythase activity: Claims 1, 6, 9 to 12, 14, 17 to 19, 23 to 27, 28 to 35 and 40 to 50 {partially}
- 8. An isolated DNA sequence coding for an enzyme involved in the carotenoid biosynthetic pathway of Phaffia rhodozyme and recombinant DNA comprising a transcription promoter and the downstream region to be expressed codes for an enzyme involved in the carotenoid biosynthesis pathway and the transformed Phaffia strains comprising said DNA, where the enzyme has phytoene desaturase activity: Claims 1, 6, 9 to 12, 14, 17 to 19, 23 to 27, 28 to 35 and 40 to 50 {partially}
- 9. An isolated DNA sequence coding for an enzyme involved in the carotenoid biosynthetic pathway of Phaffia rhodozyme and recombinant DNA comprising a transcription promoter and the downstream region to be expressed codes for an enzyme involved in the carotenoid biosynthesis pathway and the transformed Phaffia strains comprising said DNA where the enzyme has lycopene cyclase activity: Claims 1, 6, 9 to 12, 14, 17 to 19, 23 to 27, 28 to 35 and 40 to 50 (partially)
- 10. Method for the isolation of a promoter from a gene expressed in <a href="Phaffia">Phaffia</a> : Claim 52 (completely)

In. ..nation on patent family members

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